# **WEST Search History**

Hide Items Restore Clear Cancel

DATE: Saturday, September 24, 2005

Hide? Set Name Query			<b>Hit Count</b>
	DB=USPT; $PLUR=YES$ ; $OP=OR$		
	L1	6294662.pn.	1
	L2	6683156.pn.	1
	L3	6649588.pn. or 6428966.pn. or 6294662.pn. or 5916751.pn.	. 4
	L4	Tabibzadeh.in.	7

END OF SEARCH HISTORY

# **WEST Search History**

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DATE: Saturday, September 24, 2005

Hide?	<u>Set</u> Name	Query	<u>Hit</u> Count
	DB=US	SPT; PLUR=YES; OP=OR	
	L1	5874479.pn.	1
	L2	tgf\$ same (type4 or type-4 or ( type near 4))	15
	L3	(lefty1 or lefty2 or lefty-1 or lefty-2 or leftyb or lefty-b or leftya or lefty-a) and protein	6
	L4	(stra3 or stra-3) and protein	13
	DB=PC	SPB, USPT, USOC, EPAB, JPAB, DWPI, TDBD; PLUR=YES; OP=OR	
	L5	(lefty1 or lefty2 or lefty-1 or lefty-2 or leftyb or lefty-b or leftya or lefty-a) and protein	18
	L6	L5 not 13	12
	L7	L5 not 13	12
	L8	(stra3 or stra-3) and protein	96
	L9	(stra3 or stra-3) same (protein or polypeptide or peptide)	95
	L10	transforming near growth near factor near beta near 4	74
	L11	L10 not 12	74
	L12	L11 not l3	74
	L13	L12 not 15	74

END OF SEARCH HISTORY

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Trying 31060000009999...Open
DIALOG INFORMATION SERVICES
PLEASE LOGON:
 ****** HHHHHHHH SSSSSSSS? ### Status: Signing onto Dialog *******
ENTER PASSWORD:
 ****** HHHHHHHH SSSSSSS? ******
### Status: Login successfulWelcome to DIALOG
Dialog level 05.06.01D
Last logoff: 21sep05 09:05:16
Logon file405 26sep05 07:14:32
           *** ANNOUNCEMENT ***
                   ***
-- UPDATED: Important Notice to Freelance Authors--
See HELP FREELANCE for more information
NEW FILES RELEASED
***Computer and Information Systems Abstracts (File 56)
***Electronics and Communicationss Abstracts (File 57)
***Solid State and Superconductivity Abstracts (File 68)
***ANTE: Abstracts in New Technologies (File 60)
***Civil Engineering Abstracts (File 61)
***Aluminium Industry Abstracts (File 33)
***Ceramic Abstracts/World Ceramic Abstracts (File 335)
***CSA Life Sciences Abstracts (File 24)
***Corrosion Abstracts (File 46)
***Materials Business File (File 269)
***Engineered Materials Abstracts (File 293)
***CSA Aerospace & High Technology Database (File 108)
***CSA Technology Research Database (File 23)
***METADEX(r) (File 32)
***FDAnews (File 182)
***German Patents Fulltext (File 324)
RESUMED UPDATING
***Canadian Business and Current Affairs (262)
***CorpTech (559)
Chemical Structure Searching now available in Prous Science Drugs
of the Future (F453), IMS R&D Focus (F445), Beilstein Facts (F390),
and Derwent Chemistry Resource (F355).
                   ***
     >>> Enter BEGIN HOMEBASE for Dialog Announcements <<<
         of new databases, price changes, etc.
* * *
SYSTEM: HOME
Cost is in DialUnits
Menu System II: D2 version 1.7.9 term=ASCII
                     *** DIALOG HOMEBASE(SM) Main Menu ***
 Information:

    Announcements (new files, reloads, etc.)

     Database, Rates, & Command Descriptions
    Help in Choosing Databases for Your Topic
     Customer Services (telephone assistance, training, seminars, etc.)
    Product Descriptions
Connections:
```

- 6. DIALOG(R) Document Delivery
- 7. Data Star(R)
  - (c) 2003 Dialog, a Thomson business. All rights reserved.

/H = Help /L = Logoff /NOMENU = Command Mode

Enter an option number to view information or to connect to an online service. Enter a BEGIN command plus a file number to search a database (e.g., B1 for ERIC).

? b 155 medicine

Terminal set to DLINK

\*\*\* DIALOG HOMEBASE(SM) Main Menu \*\*\*

#### Information:

- Announcements (new files, reloads, etc.)
- 2. Database, Rates, & Command Descriptions
- 3. Help in Choosing Databases for Your Topic
- 4. Customer Services (telephone assistance, training, seminars, etc.)
- 5. Product Descriptions

#### Connections:

- 6. DIALOG(R) Document Delivery
- 7. Data Star(R)
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/H = Help /L = Logoff /NOMENU = Command Mode

Enter an option number to view information or to connect to an online service. Enter a BEGIN command plus a file number to search a database (e.g., B1 for ERIC).

? e transforming growth factor beta

26sep05 07:14:34 User228206 Session D2511.1 \$0.00 0.211 DialUnits FileHomeBase

- \$0.00 Estimated cost FileHomeBase
- \$0.00 Estimated cost this search
- \$0.00 Estimated total session cost 0.211 DialUnits

SYSTEM:OS - DIALOG OneSearch

File 155:MEDLINE(R) 1951-2005/Sep 26

(c) format only 2005 Dialog

File 5:Biosis Previews (R) 1969-2005/Sep W3

(c) 2005 BIOSIS

File 34:SciSearch(R) Cited Ref Sci 1990-2005/Sep W3

(c) 2005 Inst for Sci Info

File 35:Dissertation Abs Online 1861-2005/Aug

(c) 2005 ProQuest Info&Learning

File 48:SPORTDiscus 1962-2005/Apr

(c) 2005 Sport Information Resource Centre

File 65:Inside Conferences 1993-2005/Sep W3

(c) 2005 BLDSC all rts. reserv.

File 71:ELSEVIER BIOBASE 1994-2005/Sep W3

(c) 2005 Elsevier Science B.V.

File 73:EMBASE 1974-2005/Sep 26

(c) 2005 Elsevier Science B.V.

File 91:MANTIS(TM) 1880-2005/Jun

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2001 (c) Action Potential
  File 94:JICST-EPlus 1985-2005/Jul W5
         (c) 2005 Japan Science and Tech Corp (JST)
  File 98:General Sci Abs/Full-Text 1984-2004/Dec
         (c) 2005 The HW Wilson Co.
  File 135:NewsRx Weekly Reports 1995-2005/Sep W3
         (c) 2005 NewsRx
*File 135: New newsletters are now added. See Help News135 for the
complete list of newsletters.
  File 144: Pascal 1973-2005/Sep W3
         (c) 2005 INIST/CNRS
  File 149:TGG Health&Wellness DB(SM) 1976-2005/Sep W3
         (c) 2005 The Gale Group
  File 156:ToxFile 1965-2005/Sep W4
         (c) format only 2005 Dialog
  File 159:Cancerlit 1975-2002/Oct
         (c) format only 2002 Dialog
*File 159: Cancerlit is no longer updating.
Please see HELP NEWS159.
  File 162:Global Health 1983-2005/Aug
         (c) 2005 CAB International
  File 164:Allied & Complementary Medicine 1984-2005/Sep
         (c) 2005 BLHCIS
  File 172:EMBASE Alert 2005/Sep 26
         (c) 2005 Elsevier Science B.V.
  File 266:FEDRIP 2005/Jun
         Comp & dist by NTIS, Intl Copyright All Rights Res
  File 369: New Scientist 1994-2005/Jun W4
         (c) 2005 Reed Business Information Ltd.
  File 370:Science 1996-1999/Jul W3
         (c) 1999 AAAS
*File 370: This file is closed (no updates). Use File 47 for more current
information.
  File 399:CA SEARCH(R) 1967-2005/UD=14313
         (c) 2005 American Chemical Society
*File 399: Use is subject to the terms of your user/customer agreement.
Alert feature enhanced for multiple files, etc. See HELP ALERT.
  File 434:SciSearch(R) Cited Ref Sci 1974-1989/Dec
         (c) 1998 Inst for Sci Info
  File 444: New England Journal of Med. 1985-2005/Sep W2
  (c) 2005 Mass. Med. Soc. File 467:ExtraMED(tm) 2000/Dec
         (c) 2001 Informania Ltd.
*File 467: F467 no longer updates; see Help News467.
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                  TRANSFORMING GROWTH FACTOR BETA (TGF-BETA)
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                  TRANSFORMING GROWTH FACTOR BETA (TGF-BETA) AND
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                  TRANSFORMING GROWTH FACTOR BETA (TGF-BETA) REC
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39898 X DC=D24.185.348.900.720.
278463 DC=D24.35.190
DC=D24.35.540.360
DC=D24.611.350.400.800.
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         2 TRANSFORMING GROWTH FACTOR BETA (TGF-BETA) REC
         4 TRANSFORMING GROWTH FACTOR BETA (TGF-BETA) SUP
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         8 TRANSFORMING GROWTH FACTOR BETA --ADVERSE DRUG
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         66 TRANSFORMING GROWTH FACTOR BETA -- ADVERSE EFFE
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        21 TRANSFORMING GROWTH FACTOR BETA -- AGONISTS -- A
E28 2318 TRANSFORMING GROWTH FACTOR BETA --ANALYSIS --A
E29 1290 TRANSFORMING GROWTH FACTOR BETA --ANTAGONISTS
E30 4167 TRANSFORMING GROWTH FACTOR BETA --BIOSYNTHESIS
E31 1298 TRANSFORMING GROWTH FACTOR BETA --BLOOD --BL
        3 TRANSFORMING GROWTH FACTOR BETA --BUCCAL DRUG
         80 TRANSFORMING GROWTH FACTOR BETA -- CEREBROSPINA
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        2 TRANSFORMING GROWTH FACTOR BETA -- CHEMICAL SYN
       569 TRANSFORMING GROWTH FACTOR BETA -- CHEMISTRY --
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       92 TRANSFORMING GROWTH FACTOR BETA --CLASSIFICATI
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      23 TRANSFORMING GROWTH FACTOR BETA --CLINICAL TRI
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        9 TRANSFORMING GROWTH FACTOR BETA --DIAGNOSTIC U
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       15 TRANSFORMING GROWTH FACTOR BETA -- DRUG ADMINIS
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      198 TRANSFORMING GROWTH FACTOR BETA -- DRUG COMBINA
      237 TRANSFORMING GROWTH FACTOR BETA -- DRUG COMPARI
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       13 TRANSFORMING GROWTH FACTOR BETA -- DRUG CONCENT
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      102 TRANSFORMING GROWTH FACTOR BETA -- DRUG DEVELOP
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      175 TRANSFORMING GROWTH FACTOR BETA -- DRUG DOSE --
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      402 TRANSFORMING GROWTH FACTOR BETA -- DRUG EFFECTS
    105 TRANSFORMING GROWTH FACTOR BETA -- DRUG INTERAC
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       25 TRANSFORMING GROWTH FACTOR BETA -- DRUG TOXICIT
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      25 TRANSFORMING GROWTH FACTOR BETA -- DRUG TOXICIT
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      1 TRANSFORMING GROWTH FACTOR BETA --INTRAARTERIA
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         5 TRANSFORMING GROWTH FACTOR BETA --INTRAVENOUS
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       197 TRANSFORMING GROWTH FACTOR BETA -- ISOLATION AN
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         62 TRANSFORMING GROWTH FACTOR BETA --PHARMACOKINE
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         63 TRANSFORMING GROWTH FACTOR BETA -- RADIATION EF
        678 TRANSFORMING GROWTH FACTOR BETA --SECRETION --
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         4 TRANSFORMING GROWTH FACTOR BETA --STANDARDS --
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        597 TRANSFORMING GROWTH FACTOR BETA --THERAPEUTIC
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             8 TRANSFORMING GROWTH FACTOR BETA ANTIBODY
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53 TRANSFORMING GROWTH FACTOR BETA RECEPTOR --DRU
1439 TRANSFORMING GROWTH FACTOR BETA RECEPTOR --END
13 TRANSFORMING GROWTH FACTOR BETA RECEPTOR --INT
45 TRANSFORMING GROWTH FACTOR BETA RECEPTOR --PHA
        1
E28
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR --ADV
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      1439
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E34
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR --SUB
E35
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR -- TOP
E36
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR AGONI
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      Items
              RT Index-term
E37
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          1
E38
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR ALK2
E39
          1
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR ALKS
E40
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR ALK8
          1
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E41
                  TRANSFORMING GROWTH FACTOR BETA RECEPTOR ANTAG
                TRANSFORMING GROWTH FACTOR BETA RECEPTOR ASSOC TRANSFORMING GROWTH FACTOR BETA RECEPTOR BLOCK
E42
E43
E44
         1
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR EXPRE
E45
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR FAMIL
         6
E46
         20
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR I
E47
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR I KIN
         3
E48
        34
               1 TRANSFORMING GROWTH FACTOR BETA RECEPTOR II
         Enter P or PAGE for more
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Ref
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E49
          1 TRANSFORMING GROWTH FACTOR BETA RECEPTOR II GE
E50
          1 TRANSFORMING GROWTH FACTOR BETA RECEPTOR II TG
? p
            RT Index-term
Ref
      Items
E1
         1
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR II TG
E2
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR III
E3
          2
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR INHIB
E4
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR INTER
         1
E5
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR KINAS
         3
E6
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR MESSE
         1
             TRANSFORMING GROWTH FACTOR BETA RECEPTOR TGF-I

TRANSFORMING GROWTH FACTOR BETA RECEPTOR TYPE
E7
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR TGF-B
         1
E8
         25
E9
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR V
         1
E10
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR 1
         40
E11
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR 1 KIN
         1
E12
        133
               5 TRANSFORMING GROWTH FACTOR BETA RECEPTOR 2
          Enter P or PAGE for more
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      Items
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              RT Index-term
E13
         2
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR 2 -- D
E14
         65
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR 2 --E
E15
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR 2 -- I
          1
E16
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR 2 -- P
          3
E17
                   TRANSFORMING GROWTH FACTOR BETA RECEPTOR 2 MES
          1
E18
         7
                  TRANSFORMING GROWTH FACTOR BETA RECEPTOR 3
             TRANSFORMING GROWTH FACTOR BETA RECEPTORS

TRANSFORMING GROWTH FACTOR BETA REGULATORY

GROWTH FACTOR BETA REGULATORY
E19
         2
                  TRANSFORMING GROWTH FACTOR BETA RECEPTOR(S)
E20
         32
E21
         1
                   TRANSFORMING GROWTH FACTOR BETA REGULATORY ELE
E22
         1
                   TRANSFORMING GROWTH FACTOR BETA RELATED PROTEI
E23
        2
                   TRANSFORMING GROWTH FACTOR BETA RESPONSE ELEME
E24
                   TRANSFORMING GROWTH FACTOR BETA RESPONSIVE ELE
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E25
         1 TRANSFORMING GROWTH FACTOR BETA RI
E26
         1 TRANSFORMING GROWTH FACTOR BETA RII
E27
         1 TRANSFORMING GROWTH FACTOR BETA RIL
         1 TRANSFORMING GROWTH FACTOR BETA SECRETION
E28
E29
         1 TRANSFORMING GROWTH FACTOR BETA SIGNAL MEDIATO
E30
         2 TRANSFORMING GROWTH FACTOR BETA SIGNAL TRANSDU
E31
         6 TRANSFORMING GROWTH FACTOR BETA SIGNALING
E32
         5 TRANSFORMING GROWTH FACTOR BETA SIGNALING PATH
E33
         1 TRANSFORMING GROWTH FACTOR BETA SIGNALLING PAT
E34
         1 TRANSFORMING GROWTH FACTOR BETA SIMULATED CLON
E35
         1 TRANSFORMING GROWTH FACTOR BETA SOLUBLE RECEPT
E36
         4 TRANSFORMING GROWTH FACTOR BETA STIMULATED CLO
         Enter P or PAGE for more
? e transforming growth factor beta IV
Ref
     Items
             RT Index-term
E1
                 TRANSFORMING GROWTH FACTOR BETA ISOFORM OVEREX
        1
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                 TRANSFORMING GROWTH FACTOR BETA ISOFORMS
E3
                *TRANSFORMING GROWTH FACTOR BETA IV
E4
                 TRANSFORMING GROWTH FACTOR BETA L
         1
            1 TRANSFORMING GROWTH FACTOR BETA LATENCY ASSOCI
E5
E6
                 TRANSFORMING GROWTH FACTOR BETA LATENCY BINDIN
E7
                 TRANSFORMING GROWTH FACTOR BETA LIKE BINDING P
E8
                 TRANSFORMING GROWTH FACTOR BETA MASKING PROTEI
E9
         1
                 TRANSFORMING GROWTH FACTOR BETA MEMBER
E10
                 TRANSFORMING GROWTH FACTOR BETA MESSENGER RNA
                 TRANSFORMING GROWTH FACTOR BETA MONOCLONAL ANT
E11
E12
                 TRANSFORMING GROWTH FACTOR BETA MONONUCLEAR MA
         Enter P or PAGE for more
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Ref
     Items Index-term
E1
     1 TGFB3/MSX1 MARKERS
E2
         1 TGFB3VARIANT
E3
        21 *TGFB4
      1 TGFB47
E4
E5
         1 TGFB5
E6
         8 TGFC
         1 TGFCARS
E7
         1 TGFCDIA
E8
         2 TGFCYS
E9
E10
         1 TGFCYSSUP33SER
E11
         1 TGFCYS33
E12
         2 TGFCYS33SER
         Enter P or PAGE for more
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             21 'TGFB4'
     S2
? e tgfb-4
     Items Index-term
E1
        3 TGFB-3
E2
         3 TGFB-3R
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E4
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 E5
         5 TGFBA
 E6
         2 TGFBATA
 E7
         1 TGFBATTA1
         1 TGFBBA1
 E9
         1 TGFBBETA
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        1 TGFBBETA1
 E11
        1 TGFBBR2
 E12
         4 TGFBE
         Enter P or PAGE for more
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 Ref
     Items Index-term
      3 TGFBEAT
 E13
 E14
          1 TGFBEATA1
          1 TGFBEAT1
 E15
          3 TGFBEP
 E16
         1 TGFBET
 E17
 E18 14995 TGFBETA
 E19 1 TGFBETA ACTIVATED KINASE TAK1 TRANSFORMING GRO
 E20
          1 TGFBETA BETA RECEPTOR TYPE II
 E21
         1 TGFBETA EXPRESSION
 E22
         2 TGFBETA FAMILY
 E23
         1 TGFBETA FAMILY SIGNALING
         1 TGFBETA GROWTH FACTOR
 E24
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. ? p
 Ref Items Index-term
 E25
      1 TGFBETA HP75 CELLS
         1 TGFBETA INDUCED CLONE H3
E26
E27
         1 TGFBETA INHIBITOR TUMOR GROWTH FACTOR BETA INH
E28
         1 TGFBETA ISOFORMS
E29
      1 TGFBETA MODULATION
 E30
        8 TGFBETA RECEPTOR
 E31
         1 TGFBETA RECEPTOR I
E32
        1 TGFBETA RECEPTOR II
         12 TGFBETA RECEPTORS
E33
 E34
        4 TGFBETA SIGNAL TRANSDUCTION
 E35
       11 TGFBETA SIGNALING
        1 TGFBETA SIGNALING PATHWAY
E36
         Enter P or PAGE for more
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 Ref
      Items Index-term
 E37
         1 TGFBETA SIGNALING PATHWAYS
 E38
          6 TGFBETA SIGNALLING
 E39
         1 TGFBETA SIGNALS
E40
         1 TGFBETA STEROID RECEPTORS
E41
          6 TGFBETA SUPERFAMILY
E42
         1 TGFBETA TRANSFORMING GROWTH FACTOR BETA
         1 TGFBETA TRANSFORMING GROWTH FACTOR BETA TRANSF
E43
         2 TGFBETA TYPE I RECEPTOR
E44
E45
         3 TGFBETA TYPE II RECEPTOR
         1 TGFBETA TYPE II RECEPTOR PROMOTER
E46
        1 TGFBETA TYPE II RECEPTOR SIGNALING
 E47
E48
        1 TGFBETA VARIANTS
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E3

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Ref
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E49
        2 TGFBETA (TRANSFORMING GROWTH FACTOR BETA)
E50
         1 TGFBETA)
? s e48
           1 'TGFBETA VARIANTS'
? p
Ref Items Index-term
E1
         1 TGFBETA)
E2
         1 TGFBETA- 1
E3
         2 TGFBETA-DEPENDENT
E4
         1 TGFBETA-IIR
E5
         1 TGFBETA-INDEPENDENT
E6
        2 TGFBETA-INDUCED
E7
        1 TGFBETA-INDUCIBLE GENE H3 (BETAIG-H3)
        1 TGFBETA-PATHWAY
E9
        1 TGFBETA-RECEPTOR INTERACTING PROTEIN-1
E10
        3 TGFBETA-RII
E11
        1 TGFBETA-RII CELL LINE (HOMINIDAE)
E12 ·
        1 TGFBETA-SIGNALING
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Ref
    Items Index-term
     1 TGFBETA-SIGNALLING
E13
         2 TGFBETA-SUPERFAMILY
E14
E15
        1 TGFBETA-TREATED
E16
        7 TGFBETA-1
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        1 TGFBETA-1 LASER MICRODISSECTION
E18
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E19
        3 TGFBETA-2
E20
        1 TGFBETA-3
E21
         2 TGFBETA, TRANSFORMING GROWTH FACTOR BETA
E22
        1 TGFBETA, TRANSFORMING GROWTH FACTOR-BETA
E23
        2 TGFBETAA
E24
        1 TGFBETAAND
        Enter P or PAGE for more
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Ref Items Index-term
E25
        3 TGFBETABETA1
E26
        2 TGFBETAB2
E27
        1 TGFBETACALCITROIL
E28
        11 TGFBETAC10
E29
        11 TGFBETAC25
E30
        1 TGFBETAEPSILON
E31
        1 TGFBETAFNSHOWED
E32
        1 TGFBETAFNSKELETAL
        1 TGFBETAFNTREATMENT
E33
        36 TGFBETAI
E34
        1 TGFBETAIGH3
E35
       20 TGFBETAII
E36
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Ref Items Index-term
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E37
E38
        38 TGFBETAIIR
       1 TGFBETAIN
E39
      430 TGFBETAINF
E40
      12 TGFBETAINF 1
E41
       1 TGFBETAINF 3
1 TGFBETAINF 4
E42
E43
        7 TGFBETAIR
E44
        1 TGFBETAISOFORMS
E45
       16 TGFBETAL
E46
       1 TGFBETAMU
1 TGFBETAOVER
E47
E48
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Ref
     3 TGFBETAP
E49
         2 TGFBETAPRII
E50
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Ref Items Index-term
    2 TGFBETAPRII
113 TGFBETAR
E1
E2
        1 TGFBETAR II (A) 10
1 TGFBETAR II (GT) 3
E3
E4
E5
        7 TGFBETARE
        1 TGFBETARECEPTER
E6
        2 TGFBETARECEPTOR
E7
         1 TGFBETARECEPTORS
E8
        1 TGFBETARES
E9
      123 TGFBETARI
E10
       610 TGFBETARII
E11
        1 TGFBETARII FRAMESHIFT MUTATION
E12
         Enter P or PAGE for more
? p
Ref Items Index-term
E13 5 TGFBETARIIDELTAKD
        16 TGFBETARIII
E14
        7 TGFBETARS
1 TGFBETARSUB1
E15
E16
        20 TGFBETAR1
E17
       4 TGFBETAR11
30 TGFBETAR2
E18
E19
       10 TGFBETAR3
E20
       446 TGFBETAS
E21
       4 TGFBETASFR2
1 TGFBETASIGNALING
E22
E23
        1 TGFBETASL
E24
         Enter P or PAGE for more
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     Items Index-term
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     3 TGFBETASR
7 TGFBETASRII
E25
E26
E27 183 TGFBETASUB1
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1 TGFBETASUB1STIMULATED
         40 TGFBETASUB2
 E29
       1 TGFBETASUB2TGFBETASUB2
17 TGFBETASUB3
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1 TGFBETASUPERFAMILY
 E32
 E33
         1 TGFBETASUPPRESSED
 E34
         4 TGFBETASUP1
 E35
         4 TGFBETAS1
 E36
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 Ref Items Index-term
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 E37
          1 TGFBETATO
 E38
       3 TGFBETATYPE
 E39
 E40 7500 TGFBETA1
      1 TGFBETA1 - TRANSFORMING GROWTH FACTOR BETA
1 TGFBETA1 DIFFERENTIATION
1 TGFBETA1 GENE
1 TGFBETA1 MRNA
 E41
E42
 E43
 E44
         1 TGFBETA1 OVEREXPRESSION
 E45
         1 TGFBETA1 RECEPTORS
 E46
         1 TGFBETA1-ACTIVATED
 E47
         1 TGFBETA1ALPHAMP
 E48
       Enter P or PAGE for more
 ? p
 Ref Items Index-term
 E49 1 TGFBETA1ANGIOTENSINII
         1 TGFBETA1AVAILABILITY
 E50
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 Ref Items Index-term
      1 TGFBETA1AVAILABILITY
 E1
         1 TGFBETA1CDNA
 E2
        1 TGFBETA1DURING
1 TGFBETA1GENE
 E3
 E4
         1 TGFBETA1KIDNEY
 E5
 E6
         1 TGFBETA1L
 E7
         2 TGFBETA1LAP
 E8
         4 TGFBETA1MRNA
         2 TGFBETA1POSITIVE
 E9
         1 TGFBETA1PRODUCING
 E10
        11 TGFBETA1R
 E11
         1 TGFBETA1REQUIRE
 E12
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 Ref
E13 7 TGFBETA1RI
         7 TGFBETA1RII
 E14
        6 TGFBETA1SUP
 E15
         2 TGFBETA1SUPS223
 E16
         2 TGFBETA1SUPWT
 E17
         1 TGFBETA1SUP32
1 TGFBETA1TRANSGENE
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E19
       1 TGFBETA1WERE
E20
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E28

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1 TGFBETA2, IL-1
E22
         1 TGFBETA2HUMAN
E23
E24
         5 TGFBETA2R
         Enter P or PAGE for more
? p
Ref
    Items Index-term
E25
     14 TGFBETA2SUP
       934 TGFBETA3
E26
       1 TGFBETA3-ELISA
E27
        1 TGFBETA3R
E28
       1 TGFBETA3RII
E29
E30
      10 TGFBETA3SUP
      30 TGFBETA4
E31
E32
      14 TGFBETA5
       2 . TGFBETE
E33
        1 TGFBETGA1
E34
        1 TGFBGR
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E36
      375 TGFBI
        Enter P or PAGE for more
? s e31
           30 'TGFBETA4'
     S4
? p
Ref Items Index-term
     1 TGFBI (BETA IG-H3) GENE
E37
      1 TGFBI (BIGH3) GENE
5 TGFBI BIGH3 GENE
E38
E39
E40
        15 TGFBI GENE
       1 TGFBI TRANSFORMING GROWTH FACTOR-BETA-1
E41
        1 TGFBI(BIGH3) GENE
E42
        1 TGFBI-INDUCED
E43
        2 TGFBIGENE
E44
E45
        9 TGFBII
        1 TGFBIII
E46
       6 TGFBIIIR
E47
       44 TGFBIIR
E48
        Enter P or PAGE for more
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     Items Index-term
Ref
E49
        1 TGFBIIR GENE
E50
        1 TGFBINDUCES
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Ref Items Index-term
E1
        1 TGFBINDUCES
E2
        7 TGFBINF
E3
        2 TGFBIN3
E4
        19 TGFBIP
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        7 TGFBKM2
E6
        2 TGFBKM2SUP129
E7
        2 TGFBKM2129
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       7 TGFBM1
E9
       6 TGFBM3
E10
      25 TGFBP
E11
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E21 1402 TGFBETA2

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Ref Items Index-term
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E13
E14
         1 TGFBP1
E15
        1 TGFBP3
E16
        1 TGFBP4
E17
        1 TGFBP5
        27 TGFBR
E18
       1 TGFBR-SMAD
E19
        1 TGFBR-SMAD SUPERFAMILY
E20
E21
        11 TGFBRAP1
        11 TGFBRAP1 PROTEIN, HUMAN
E22
E23
       21 TGFBRI
      111 TGFBRII
E24
        Enter P or PAGE for more
? s e16
            1 'TGFBP4'
     S5
? p
    Items Index-term
Ref
E25
        1 TGFBRII GENE
         1 TGFBRII GENE TRANSFORMING GROWTH FACTOR-BETA R
E26
E27
        3 TGFBRL
        2 TGFBRLOXP
E28
        1 TGFBRMRNA
E29
       6 TGFBRR1
E30
       6 TGFBRR2
E31
E32
        1 TGFBRS
        2 TGFBRTA
E33
E34
       163 TGFBR1
E35
        1 TGFBR1 GENE
E36
        1 TGFBR1 KINASE
         Enter P or PAGE for more
? p
Ref Items Index-term
E37
        1 TGFBR1 TRANSFORMING GROWTH FACTOR, BETA RECEPT
E38
         4 TGFBR1-ASTERISK-6A
E39
        1 TGFBR1-6A ALLELE
E40
        2 TGFBR16A
E41
       463 TGFBR2
E42
        9 TGFBR2 GENE
        1 TGFBR2 GENE TYPE II TRANSFORMING GROWTH FACTOR
        13 TGFBR2 PROTEIN, HUMAN
E45
        2 TGFBR2-RESTORED
E46
        1 TGFBR2FLOXE2
E47
        2 TGFBR2FLX
E48
        4 TGFBR2FSPKO
         Enter P or PAGE for more
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Ref
     Items Index-term
E49
      1 TGFBR2HEPKO
E50
        1 TGFBR2L
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? e lefty 1
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     Items Index-term
E1
         1 LEFTY SIGNALING
E2
         1 LEFTY VENTRICULAR REMODELING
         0 *LEFTY 1
E3
         1 LEFTY 1 PROTEIN
E4
         1 LEFTY- 1
E5
         4 LEFTY-A
E6
E7 .
        1 LEFTY-A ENDOMETRIAL BLEEDING-ASSOCIATED FACTOR
        1 LEFTY-A MRNA
E8
        1 LEFTY-A/ENDOMETRIAL BLEEDING-ASSOCIATED FACTOR
E9
        2 LEFTY-B
E10
        1 LEFTY-DEPENDENT
E11
E12
         1 LEFTY-GENE RELATED PRODUCT
         Enter P or PAGE for more
? s e4-e10
              1 LEFTY 1 PROTEIN
              1 LEFTY- 1
              4 LEFTY-A
              1 LEFTY-A ENDOMETRIAL BLEEDING-ASSOCIATED FACTOR
              1 LEFTY-A MRNA
              1 LEFTY-A/ENDOMETRIAL BLEEDING-ASSOCIATED FACTOR
              2 LEFTY-B
     S6
             8 E4-E10
? p
Ref
    Items Index-term
        1 LEFTY-RELATED
E13
        1 LEFTY-SIGNALING
E14
        13 LEFTY-1
E15
        5 LEFTY-1 GENE
E16
        1 LEFTY-1 PROTEIN
E17
        12 LEFTY-2
E18
        3 LEFTY-2 GENE
E19
E20
        1 LEFTY-2 ISOLATION
E21
        1 LEFTY-2 PROTEIN
E22
        34 LEFTYA
E23
        1 LEFTYA GENE
        1 LEFTYA PROTEIN
E24
         Enter P or PAGE for more
? s e15 or e17 or e18 or e20 or e21 or e22 or e24
             13 LEFTY-1
              1 LEFTY-1 PROTEIN
             12 LEFTY-2
              1 LEFTY-2 ISOLATION
              1 LEFTY-2 PROTEIN
             34 LEFTYA
             1 LEFTYA PROTEIN
     S7
             57 'LEFTY-1' OR 'LEFTY-1 PROTEIN' OR 'LEFTY-2' OR 'LEFTY-2
                 ISOLATION' OR 'LEFTY-2 PROTEIN' OR 'LEFTYA' OR 'LEFTYA
               PROTEIN
? p
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     Items Index-term
E25
        2 LEFTYB
E26
        1 LEFTYB PROTEIN
E27
        5 LEFTYL
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1 LEFTYRAY

E28

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2 LEFTYS
E29
      204 LEFTY1
E30
       3 LEFTY1 GENE
E31
E32
         1 LEFTY1 PROTEIN
E33
         1 LEFTY1GENE
        6 LEFTY1LEFTY2
E34
E35
         1 LEFTY1LEFTY2 TUBULIN MUTANT
E36
         4 LEFTY1SUP
         Enter P or PAGE for more
? s e25-e26 or e30 or e32 or e34 or e35 or e36
              2 LEFTYB
             1 LEFTYB PROTEIN
            204 LEFTY1
              1 LEFTY1 PROTEIN
              6 LEFTY1LEFTY2
             1 LEFTY1LEFTY2 TUBULIN MUTANT
             4 LEFTY1SUP
            206 E25-E26 OR E30 OR E32 OR E34 OR E35 OR E36
     S8
? p
Ref
     Items Index-term
     196 LEFTY2
E37
        3 LEFTY2 GENE
E38
         1 LEFTY2 PROTEIN
E39
E40
         1 LEFTY2GENE
E41
         1 LEFTY3
E42
        1 LEFTY69
E43
        4 LEFT1
E44
        1 LEFU
E45
        91 LEFUA
       1 LEFUA (OSTEICHTHYES)
E46
        6 LEFUA COSTATA
E47
E48
        6 LEFUA COSTATA (OSTEICHTHYES)
         Enter P or PAGE for more
? s e37 or e39
            196 LEFTY2
            1 LEFTY2 PROTEIN
            196 'LEFTY2' OR 'LEFTY2 PROTEIN'
? p
Ref
     Items Index-term
        1 LEFUA COSTATA ECHIGONIA
E49
         1 LEFUA COSTATA MITOCHONDRIAL D-LOOP REGION GENE
E50
? e lefty a
     Items Index-term
Ref
       929 LEFTY
E1
         1 *LEFTY A
E2
E3
         6 LEFTY GENE
E4
         1 LEFTY KREH'S ULTIMATE GUIDE TO FLY FISHING (BO
E5
         2 LEFTY MUTANTS
E6
        1 LEFTY POPULATION
E7 '
       155 LEFTY PROTEIN
        1 LEFTY PROTEIN FAMILY
        4 LEFTY PROTEINS
E9
       1 LEFTY SECRETED FACTORS
E10
E11
       1 LEFTY SEOUENCE
   1 LEFTY SIGNALING
E12
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Enter P or PAGE for more
? s e2 or e7 or e8 or e9
              1 LEFTY A
            155 LEFTY PROTEIN
              1 LEFTY PROTEIN FAMILY
              4 LEFTY PROTEINS
            161 'LEFTY A' OR 'LEFTY PROTEIN' OR 'LEFTY PROTEIN FAMILY' OR
    S10
                 'LEFTY PROTEINS'
? p
Ref
     Items Index-term
        1 LEFTY VENTRICULAR REMODELING
E13
         1 LEFTY 1 PROTEIN
E14
         1 LEFTY- 1
E15
         4 LEFTY-A
E16
E17
         1 LEFTY-A ENDOMETRIAL BLEEDING-ASSOCIATED FACTOR
         1 LEFTY-A MRNA
E18
         1 LEFTY-A/ENDOMETRIAL BLEEDING-ASSOCIATED FACTOR
E19
         2 LEFTY-B
E20
        1 LEFTY-DEPENDENT
E21
         1 LEFTY-GENE RELATED PRODUCT
E22
        1 LEFTY-RELATED
E23
         1 LEFTY-SIGNALING
E24
         Enter P or PAGE for more
? s e14 or e15 or e16 or e17 or e18 or e19 or e20
              1 LEFTY 1 PROTEIN
              1 LEFTY- 1
              4 LEFTY-A
              1 LEFTY-A ENDOMETRIAL BLEEDING-ASSOCIATED FACTOR
              1 LEFTY-A MRNA
              1 LEFTY-A/ENDOMETRIAL BLEEDING-ASSOCIATED FACTOR
              2 LEFTY-B
              8 'LEFTY 1 PROTEIN' OR 'LEFTY- 1' OR 'LEFTY-A' OR 'LEFTY-A
    S11
                 ENDOMETRIAL BLEEDING-ASSOCIATED FACTOR' OR 'LEFTY-A MRNA'
                 OR 'LEFTY-A/ENDOMETRIAL BLEEDING-ASSOCIATED FACTOR' OR
                 'LEFTY-B'
? ds
Set
              Description
       Items
              'TRANSFORMING GROWTH FACTOR BETA ISOFORMS'
S1
          3
S2
              'TGFB4'
          21
S3
          1
              'TGFBETA VARIANTS'
S4
          30
              'TGFBETA4'
              'TGFBP4'
S5
           1
               E4-E10
S6
           8
              'LEFTY-1' OR 'LEFTY-1 PROTEIN' OR 'LEFTY-2' OR 'LEFTY-2 IS-
S7
          57
            OLATION' OR 'LEFTY-2 PROTEIN' OR 'LEFTYA' OR 'LEFTYA PROTEIN'
         206 E25-E26 OR E30 OR E32 OR E34 OR E35 OR E36
S8
               'LEFTY2' OR 'LEFTY2 PROTEIN'
S9
         196
              'LEFTY A' OR 'LEFTY PROTEIN' OR 'LEFTY PROTEIN FAMILY' OR -
S10
         161
            'LEFTY PROTEINS'
               'LEFTY 1 PROTEIN' OR 'LEFTY- 1' OR 'LEFTY-A' OR 'LEFTY-A E-
S11
            NDOMETRIAL BLEEDING-ASSOCIATED FACTOR' OR 'LEFTY-A MRNA' OR '-
            LEFTY-A/ENDOMETRIAL BLEEDING-ASSOCIATED FACTOR' OR 'LEFTY-B'
? s s1 or s2 or s3 or s4 or s5 or s6 or s7 or s8 or s9 or s10 or s11
              3 S1
             21 S2
              1 S3
             30 S4
```

```
1 S5
               8
                 S6
              57
                 S7
             206 S8
             196 S9
             161 S10
                 S11
               8
     S12
             531 S1 OR S2 OR S3 OR S4 OR S5 OR S6 OR S7 OR S8 OR S9 OR S10
                  OR S11
? s s12/1996:2005
Processing
Processed 10 of 26 files ...
>>>One or more prefixes are unsupported
>>> or undefined in one or more files.
>>>Year ranges not supported in one or more files
Processing
Completed processing all files
             531 S12
        49727561 PY=1996 : PY=2005
     S13
             525 S12/1996:2005
? s s12 not s13
             531 S12
             525 S13
               6 S12 NOT S13
     S14
? rd
...completed examining records
     S15
               4 RD (unique items)
? t s15/6/all
            (Item 1 from file: 71)
15/6/1
00304408
                95125590
The immunomodulatory diversity of the proteins of the transforming growth
  factor beta (TGFbetaP) family
PUBLICATION DATE: 19950000
 15/6/2
            (Item 2 from file: 71)
00290512
                95107663
Expression of transforming growth factor beta in the embryonic avian lens
  coincides with the presence of mitochondria
PUBLICATION DATE: 19950000
 15/6/3
            (Item 1 from file: 144)
            PASCAL No.: 92-0692127
  10488633
  Localization of the human insulin-like growth-factor-binding protein 4
gene to chromosomal region 17q12-21.1
  1992
            (Item 1 from file: 399)
DIALOG(R) File 399:(c) 2005 American Chemical Society. All rts. reserv.
  Complementary deoxyribonucleic acid cloning of a messenger ribonucleic
acid encoding transforming growth factor .beta. 4 from chicken embryo
chondrocytes
? t s15/9/1 2 3
15/9/1
            (Item 1 from file: 71)
```

DIALOG(R) File 71:ELSEVIER BIOBASE (c) 2005 Elsevier Science B.V. All rts. reserv.

00304408 95125590

The immunomodulatory diversity of the proteins of the transforming growth factor beta (TGFbetaP) family

Wieczorek Z.; Sion J.; Kluczyk A.; Zbozien R.; Stafanowicz P.; Siemion I.Z. ADDRESS: Z. Wieczorek, L. Hirschfeld Inst Immun/Exp Therapy, Polish Academy

of Sciences, Czerska 12, 53-114 Wrocław, Poland Journal: International Journal of Peptide and Protein Research, 46/2 (113-118), 1995, Denmark

PUBLICATION DATE: 19950000

CODEN: IJPPC ISSN: 0367-8377

DOCUMENT TYPE: Article

LANGUAGES: English SUMMARY LANGUAGES: English

The examination of immunomodulatory properties of oligopeptides derived from two exposed loops (containing thymopentin-like and tuftsin-like sequences, respectively) of the proteins belonging to TGFbeta family suggests that the particular species of the TGFbeta family should differ distinctly in their influence on the immune response. According to our results obtained from three TGFbeta species of mammals, TGFbeta 2 should be a strong immunosuppressor, whereas for TGFbeta 3 the immunostimulative potency is more probable. TGFbeta 1 species would possess both immunosuppressive and immunostimulative potency, residing in two different loops of the protein. The results obtained also suggest that chicken TGFbeta4 should be associated with immunostimulative effects and xenopus TGFbeta5 with immunosuppressive ones.

#### **DESCRIPTORS:**

Peptide immunomodulators; Thymopentin-like sequences; Transforming growth factor beta; Tuftsin-like sequences

CLASSIFICATION CODE AND DESCRIPTION:

89.2.4.5 - CELL AND DEVELOPMENTAL BIOLOGY / CELL GROWTH AND DIVISION /
 Growth Factors and Inhibitors / Transforming growth factors (TGF)
86.5.4.6 - IMMUNOLOGY AND INFECTIOUS DISEASES / HUMORAL MEDIATORS OF IMMUNE
 RESPONSE / Other Factors / Haematopoietic growth factors (CSF)

15/9/2 (Item 2 from file: 71)
DIALOG(R)File 71:ELSEVIER BIOBASE
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00290512 95107663

Expression of transforming growth factor beta in the embryonic avian lens coincides with the presence of mitochondria

Potts J.D.; Bassnett S.; Beebe D.C.

ADDRESS: J.D. Potts, Department of Anatomy/Cell Biology, Uniformed Svcs.
Univ. of Health Sci., 4301 Jones Bridge Road, Bethesda, MD
20814-4799, United States

Journal: Developmental Dynamics, 203/3 (317-323), 1995, United States PUBLICATION DATE: 19950000

CODEN: DEDYE ISSN: 1058-8388

DOCUMENT TYPE: Article

LANGUAGES: English SUMMARY LANGUAGES: English

During their maturation, lens cells lose all membrane bound organelles, including mitochondria. In chicken embryos this process begins in the

central lens fibers beginning around embryonic day 12 (E12). Transforming growth factor beta (TGFbeta) is a multipotent growth modulator thought to play a role in numerous developmental processes. TGFbeta1 has been localized to mitochondria in rat liver cells and muscle cells. In the present study, we examined the expression of TGFbeta isoform mRNAs and proteins during chicken embryonic lens development. PCR analysis demonstrated TGFbeta2 and TGFbeta3 transcripts in the lens epithelium and fibers throughout pre- and post-hatching development. TGFbeta isoforms were detected throughout the lens epithelium and fibers early in development (E6). However by E19, the distribution of TGFbeta2 and TGFbeta3 transcripts and proteins coincided with regions of the lens that contained mitochondria. In addition, intense TGFbeta staining was observed in the basal portions of the equatorial epithelial cells, a region with abundant mitochondria. Transcripts for TGFbeta1 and TGFbeta4 were not detected in any tissue or time frame examined. Similarly, no immunostaining for TGFbetal was observed.

#### DESCRIPTORS:

TGFbeta; Chicken lens; Lens differentiation; Mitochondria

CLASSIFICATION CODE AND DESCRIPTION:

89.8.9.7 - CELL AND DEVELOPMENTAL BIOLOGY / DEVELOPMENT (BY TISSUE AND ORGAN SYSTEMS) / Nervous System / Sensory - visual

89.2.4.5 - CELL AND DEVELOPMENTAL BIOLOGY / CELL GROWTH AND DIVISION / Growth Factors and Inhibitors / Transforming growth factors (TGF)

89.1.8.7 - CELL AND DEVELOPMENTAL BIOLOGY / MEMBRANES AND CELL TRANSPORT / Cytoplasmic Membranes / Mitochondria

15/9/3 (Item 1 from file: 144)
DIALOG(R)File 144:Pascal
(c) 2005 INIST/CNRS. All rts. reserv.

10488633 PASCAL No.: 92-0692127

Localization of the human insulin-like growth-factor-binding protein 4 gene to chromosomal region 17q12-21.1

BAJALICA S; ALLANDER S V; EHRENBORG E; BRONDUM-NIELSEN K; LUTHMAN H; LARSSON C

Karolinska hosp., dep. clinical genetics, 10401 Stockholm, Sweden Journal: Human genetics, 1992, 89 (2) 234-236

ISSN: 0340-6717 CODEN: HUGEDQ Availability: INIST-2672; 354000028118920170

No. of Refs.: 13 ref.

Document Type: P (Serial) ; A (Analytic)

Country of Publication: Federal Republic of Germany

Language: English

Insulin-like growth-factor-binding proteins (IGFBPs) constitute a family of structurally related proteins that specifically bind insulin-like growth factors and modulate their functions. In this study, the chromosomal gene encoding IGFBP4, i.e. determined localization was for the inhibitory-IGFBP. A polymerase chain reaction (PCR) fragment corresponding to the previously published cDNA sequence was used to isolate overlapping clones. By fluorescent in situ hybridization to metaphase cosmid TGFBP4 chromosomes, the gene was then localized to chromosomal region 17q21-21.1

English Descriptors: E17-Chromosome; Binding protein; Genetic mapping;
Human; Somatomedin C; Insulin like growth factor 2; Polymerase chain
reaction; Molecular hybridization; In situ; Molecular biology; Genetics;
Exploration

French Descriptors: Chromosome E17; Proteine liaison; Carte genetique; Homme; Somatomedine C; Facteur croissance IGF2; Reaction chaine polymerase; Hybridation moleculaire; In situ; Biologie moleculaire; Genetique; Exploration Classification Codes: 002A07C03 ? t s15/9/4 15/9/4 (Item 1 from file: 399) DIALOG(R) File 399:CA SEARCH(R) (c) 2005 American Chemical Society. All rts. reserv. **JOURNAL** 111091468 CA: 111(11)91468h Complementary deoxyribonucleic acid cloning of a messenger ribonucleic acid encoding transforming growth factor .beta. 4 from chicken embryo chondrocytes AUTHOR(S): Jakowlew, Sonia B.; Dillard, Pamela J.; Sporn, Michael B.; Roberts, Anita B. LOCATION: Lab. Chemoprev., Natl. Cancer Inst., Bethesda, MD, 20892, USA JOURNAL: Mol. Endocrinol. DATE: 1988 VOLUME: 2 NUMBER: 12 PAGES: 1186-95 CODEN: MOENEN ISSN: 0888-8809 LANGUAGE: English SECTION: CA203003 Biochemical Genetics CA202XXX Mammalian Hormones CA213XXX Mammalian Biochemistry IDENTIFIERS: cDNA cloning sequencing chicken TGFbeta4, transforming growth factor chicken gene sequence **DESCRIPTORS:** Gene and Genetic element, animal... for transforming growth factor .beta.4, of chicken, nucleotide and encoded peptide sequences of Animal growth regulators, .beta.4-transforming growth factors... gene for, of chicken, nucleotide and encoded peptide sequences of Deoxyribonucleic acid sequences, beta.4-transforming growth factor-specifying... of chicken, multiple clones, complete Protein sequences... of transforming growth factor .beta.4 and precursor, multiple clones, complete Molecular cloning... of transforming growth factor .beta.4 cDNA, of chicken Chicken... transforming growth factor .beta.4 gene of, nucleotide and encoded peptide sequences of CAS REGISTRY NUMBERS: 122319-69-1 122319-70-4 amino acid sequence of 122318-73-4 nucleotide sequence of ? logoff hold 26sep05 07:24:19 User228206 Session D2511.2 \$2.83 0.832 DialUnits File155 \$2.83 Estimated cost File155 0.907 DialUnits File5 \$5.35 \$5.35 Estimated cost File5 1.323 DialUnits File34 \$29.30 \$29.30 Estimated cost File34 \$0.87 0.211 DialUnits File35 Estimated cost File35 \$1.06 0.198 DialUnits File48 \$1.06 Estimated cost File48

1.378 DialUnits File65

\$5.17

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$5.17 Estimated cost File65
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         $5.25
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           $4.00 2 Type(s) in Format 9
        $4.00 4 Types
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                0.300 DialUnits File156
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           $0.55 1 Type(s) in Format 6
           $2.75 1 Type(s) in Format 9
        $3.30 2 Types
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 $2.57 Estimated cost File434
                 0.068 DialUnits File444
        $0.33
 $0.33 Estimated cost File444
                0.096 DialUnits File467
        $0.61
 $0.61 Estimated cost File467
        OneSearch, 26 files, 10.996 DialUnits FileOS
 $2.66 TELNET
$101.84 Estimated cost this search
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\$101.84 Estimated total session cost 11.207 DialUnits

Logoff: level 0 You are now logged off

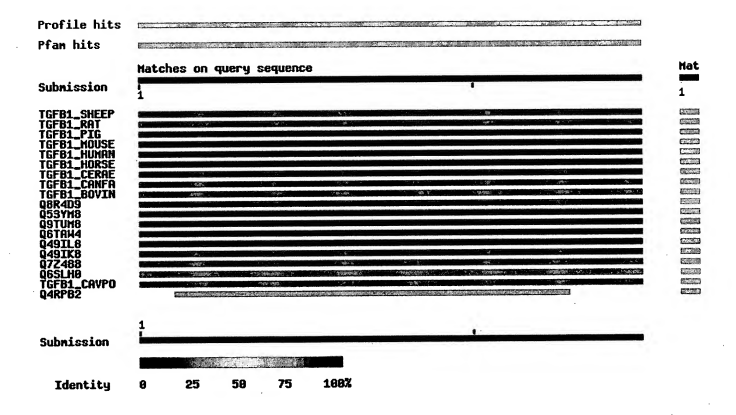
ExPASy Home page Site Map Search ExPASy Contact us Proteomics tools Swiss-Prot
Search Swiss-Prot/TrEMBL for tgfb 1 @ Clear
Search Swiss-Prot/TrEMBL of for tgfb 1 Go Clear  Welcome to the SIB BLAST Network Service
Welcome to the SIB BLAST Network Service
welcome to the SIB BLASI Network Service
If results of this search are reported or published, please mention that the computation was performed at the SIB using the BLAST network service.  The SIB BLAST network service uses a server developed at SIB and the NCBI BLAST 2 software.
In case of problems, please read the <u>online BLAST help</u> .  If your question is not covered, please contact < <u>helpdesk@expasy.org</u> >.
NCBI BLAST program reference [PMID: 9254694]: Altschul S.F., Madden T.L., Schäffer A.A., Zhang J., Zhang Z., Miller W., Lipman D.J. Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. Nucleic Acids Res. 25:3389-3402(1997).
Query: 15 AA  Date run: 2005-09-24 15:42:21 UTC+0100 on sib-gml.unil.ch  Program: NCBI BLASTP 1.5.4-Paracel [2003-06-05]  Database: EXPASY/UniProtKB  2,312,053 sequences; 758,486,757 total letters  UniProt Knowledgebase Release 6.0 consists of:  UniProtKB/Swiss-Prot Release 48.0 of 13-Sep-2005: 194317 entries  UniProtKB/TrEMBL Release 31.0 of 13-Sep-2005: 2105517 entries
Taxonomic view NiceBlast view Printable view
List of potentially matching sequences
Send selected sequences to Clustal W (multiple alignment)  Select up to Submit Query  Include query sequence
Db AC Description Score E-value
sp P50414 TGFB1 SHEEP Transforming growth factor beta 1 precurso 53 6e-07
sp P17246 TGFB1 RAT Transforming growth factor beta 1 precursor 53 6e-07
sp P07200 TGFB1 PIG Transforming growth factor beta 1 precursor 53 6e-07
sp P04202 TGFB1_MOUSE Transforming growth factor beta 1 precurso 53 6e-07
sp P01137 TGFB1_HUMAN Transforming growth factor beta 1 precurso 53 6e-07
sp 019011 TGFB1 HORSE Transforming growth factor beta 1 precurso 53 6e-07
sp P09533 TGFB1_CERAE Transforming growth factor beta 1 precurso 53 6e-07
sp P54831 TGFB1_CANFA Transforming growth factor beta 1 precurso 53 6e-07
sp P18341 TGFB1_BOVIN Transforming growth factor beta 1 precurso 53 6e-07

```
tr Q8R4D9 _SIGHI Transforming growth factor beta-1 protein (Fragm... 53 6e-07 tr Q53YM8 _RAT TGF beta 1 [Rattus norvegicus (Rat)] 53 6e-07 tr Q9TUM8 _HORSE Transforming growth factor beta 1 [TGFb1] [Equus... 53 6e-07 tr Q6TAW4 _FELCA TGF beta (Fragment) [Felis silvestris catus (Cat)] 53 6e-07 tr Q49IL8 _HUMAN Transforming growth factor beta 1 precursor (Fra... 53 6e-07 tr Q49IK8 _HUMAN Transforming growth factor beta 1 precursor (Fra... 53 6e-07 tr Q7Z488 _HUMAN Transforming growth factor beta 1 precursor (Fra... 53 6e-07 tr Q6SLHO _PERMA Transforming growth factor beta 1 (Fragment) [Tg... 50 5e-06 sp Q9Z1Y6 TGFB1_CAVPO Transforming growth factor beta 1 precurso... 50 7e-06 tr Q4RPB2 _TETNG Chromosome 1 SCAF15008, whole genome shotgun seq... 32 2.1
```

### Graphical overview of the alignments

Click here to resubmit your query after masking regions matching PROSITE profiles or Pfam HMMs

( Help) (use <a href="ScanProsite">ScanProsite</a> for more details about PROSITE matches)



### Alignments

```
sp P50414 Transforming growth factor beta 1 precursor (TGF-beta 1) 390
AA
[Ovis aries (Sheep)] align

Score = 53.2 bits (118), Expect = 6e-07
Identities = 15/15 (100%), Positives = 15/15 (100%)
```

Query: 1

PEADYYAKEVTRVLM 15

PEADYYAKEVTRVLM Sbjct: 99 PEADYYAKEVTRVLM 113 **sp** P17246 Transforming growth factor beta 1 precursor (TGF-beta 1) 390 TGFB1 RAT [Tgfb1] AA [Rattus norvegicus (Rat)] align Score = 53.2 bits (118), Expect = 6e-07Identities = 15/15 (100%), Positives = 15/15 (100%) Query: 1 PEADYYAKEVTRVLM 15 PEADYYAKEVTRVLM Sbjct: 99 PEADYYAKEVTRVLM 113 **sp** P07200 390 Transforming growth factor beta 1 precursor (TGF-beta 1) TGFB1 PIG [TGFB1] AΑ [Sus scrofa (Pig)] align Score = 53.2 bits (118), Expect = 6e-07Identities = 15/15 (100%), Positives = 15/15 (100%) Query: 1 PEADYYAKEVTRVLM 15 PEADYYAKEVTRVLM Sbjct: 99 PEADYYAKEVTRVLM 113 **sp** P04202 Transforming growth factor beta 1 precursor (TGF-beta 1) 390 TGFB1 MOUSE [Tgfb1] AA [Mus musculus (Mouse)] align Score = 53.2 bits (118), Expect = 6e-07Identities = 15/15 (100%), Positives = 15/15 (100%) PEADYYAKEVTRVLM 15 Query: 1 PEADYYAKEVTRVLM Sbjct: 99 PEADYYAKEVTRVLM 113 **sp** P01137 Transforming growth factor beta 1 precursor (TGF-beta 1) 390 TGFB1 HUMAN [TGFB1] AΑ [Homo sapiens (Human)] align Score = 53.2 bits (118), Expect = 6e-07 Identities = 15/15 (100%), Positives = 15/15 (100%) Query: 1 PEADYYAKEVTRVLM 15 PEADYYAKEVTRVLM Sbjct: 99 PEADYYAKEVTRVLM 113

sp 019011 Transforming growth factor beta 1 precursor (TGF-beta 1) 390
TGFB1\_HORSE [TGFB1] AA
[Equus caballus (Horse)] align

Score = 53.2 bits (118), Expect = 6e-07
Identities = 15/15 (100%), Positives = 15/15 (100%)

Query: 1 PEADYYAKEVTRVLM 15
. PEADYYAKEVTRVLM
Sbjct: 99 PEADYYAKEVTRVLM 113

sp P09533 Transforming growth factor beta 1 precursor (TGF-beta 1) 390 TGFB1\_CERAE [TGFB1] AA

[Cercopithecus aethiops (Green monkey) (Grivet)] align

Score = 53.2 bits (118), Expect = 6e-07
Identities = 15/15 (100%), Positives = 15/15 (100%)

Query: 1 PEADYYAKEVTRVLM 15
PEADYYAKEVTRVLM
Sbjct: 99 PEADYYAKEVTRVLM 113

sp P54831 Transforming growth factor beta 1 precursor (TGF-beta 1) 390
TGFB1\_CANFA [TGFB1] AA
[Canis familiaris (Dog)] align

Score = 53.2 bits (118), Expect = 6e-07
Identities = 15/15 (100%), Positives = 15/15 (100%)

Query: 1 PEADYYAKEVTRVLM 15
PEADYYAKEVTRVLM
Sbjct: 99 PEADYYAKEVTRVLM 113

sp P18341 Transforming growth factor beta 1 precursor (TGF-beta 1) 315 AA TGFB1\_BOVIN (Fragment) [TGFB1] [Bos taurus (Bovine)] align

Score = 53.2 bits (118), Expect = 6e-07 Identities = 15/15 (100%), Positives = 15/15 (100%)

Query: 1 PEADYYAKEVTRVLM 15
PEADYYAKEVTRVLM 5
Sbjct: 24 PEADYYAKEVTRVLM 38

Transforming growth factor beta-1 protein (Fragment) 368 tr Q8R4D9 Q8R4D9 SIGHI [Tgfb1] AΑ align [Sigmodon hispidus (Hispid cotton rat)] Score = 53.2 bits (118), Expect = 6e-07 Identities = 15/15 (100%), Positives = 15/15 (100%) Query: 1 PEADYYAKEVTRVLM 15 PEADYYAKEVTRVLM Sbjct: 77 PEADYYAKEVTRVLM 91 tr Q53YM8 TGF beta 1 [Rattus norvegicus (Rat)] 390 AA Q53YM8 RAT align Score = 53.2 bits (118), Expect = 6e-07 Identities = 15/15 (100%), Positives = 15/15 (100%) Query: 1 PEADYYAKEVTRVLM 15 PEADYYAKEVTRVLM Sbjct: 99 PEADYYAKEVTRVLM 113 Transforming growth factor beta 1 [TGFb1] [Equus 390 tr Q9TUM8 AΑ Q9TUM8\_HORSE caballus (Horse)] align Score = 53.2 bits (118), Expect = 6e-07 Identities = 15/15 (100%), Positives = 15/15 (100%) PEADYYAKEVTRVLM 15 Query: 1 PEADYYAKEVTRVLM Sbjct: 99 PEADYYAKEVTRVLM 113 tr Q6TAW4 TGF beta (Fragment) [Felis silvestris catus (Cat)] 226 AA Q6TAW4 FELCA align Score = 53.2 bits (118), Expect = 6e-07 Identities = 15/15 (100%), Positives = 15/15 (100%) Query: 1 PEADYYAKEVTRVLM 15 PEADYYAKEVTRVLM Sbjct: 50 PEADYYAKEVTRVLM 64 Transforming growth factor beta 1 precursor (Fragment) 118 tr Q49IL8 Q49IL8 HUMAN [TGFB1] AΑ

[Homo sapiens (Human)]

align

Score = 53.2 bits (118), Expect = 6e-07Identities = 15/15 (100%), Positives = 15/15 (100%) Query: 1 PEADYYAKEVTRVLM 15 PEADYYAKEVTRVLM Sbjct: 99 PEADYYAKEVTRVLM 113 tr Q49IK8 Transforming growth factor beta 1 precursor (Fragment) Q491K8 HUMAN [TGFB1] AΑ [Homo sapiens (Human)] align Score = 53.2 bits (118), Expect = 6e-07 Identities = 15/15 (100%), Positives = 15/15 (100%) Query: 1 PEADYYAKEVTRVLM 15 PEADYYAKEVTRVLM Sbjct: 99 PEADYYAKEVTRVLM 113 tr Q7Z488 Transforming growth factor beta 1 precursor (Fragment) 118 Q7Z488 HUMAN [TGFB1] AΑ [Homo sapiens (Human)] align Score = 53.2 bits (118), Expect = 6e-07 Identities = 15/15 (100%), Positives = 15/15 (100%) Query: 1 PEADYYAKEVTRVLM 15 PEADYYAKEVTRVLM Sbjct: 99 PEADYYAKEVTRVLM 113 Transforming growth factor beta 1 (Fragment) [Tgfb1] tr Q6SLH0 249 Q6SLH0 PERMA [Peromyscus AA align maniculatus (Deer mouse)] Score = 50.3 bits (111), Expect = 5e-06Identities = 14/15 (93%), Positives = 14/15 (93%) Query: 1 PEADYYAKEVTRVLM 15 PE DYYAKEVTRVLM Sbjct: 65 PETDYYAKEVTRVLM 79 **sp** Q9Z1Y6 Transforming growth factor beta 1 precursor (TGF-beta 1) 390 TGFB1\_CAVPO [TGFB1] AA [Cavia porcellus (Guinea pig)] align Score = 49.8 bits (110), Expect = 7e-06Identities = 14/15 (93%), Positives = 14/15 (93%)

Query: 1

PEADYYAKEVTRVLM 15

```
PE DYYAKEVTRVLM
 Sbjct: 99 PEPDYYAKEVTRVLM 113
                  Chromosome 1 SCAF15008, whole genome shotgun sequence.
                                                                             454
 tr Q4RPB2
     Q4RPB2 TETNG (Fragment)
                                                                             AΑ
                  [GSTENG00031190001] [Tetraodon nigroviridis (Green
                                                                             align
                  puffer)]
  Score = 31.6 bits (67), Expect = 2.1
  Identities = 9/12 (75%), Positives = 10/12 (83%)
 Query: 2
           EADYYAKEVTRV 13
            E DYYAKEV R+
 Sbjct: 96 EEDYYAKEVQRI 107
Database: EXPASY/UniProtKB
    Posted date: Sep 12, 2005 12:34 PM
 Number of letters in database: 758,486,757
 Number of sequences in database: 2,312,053
Lambda
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           0.280
                      1.74
Gapped
Lambda
          K
                 Η
   0.294
          0.110
                     0.610
Matrix: PAM30
Gap Penalties: Existence: 9, Extension: 1
Number of HSP's successfully gapped in prelim test: 0
length of query: 15
length of database: 758,486,757
effective HSP length: 6
effective length of query: 9
effective length of database: 744,614,439
effective search space: 6701529951
effective search space used: 6701529951
T: 16
A: 40
X1: 15 (7.1 bits)
X2: 35 (14.8 bits)
X3: 58 (24.6 bits)
S1: 42 (21.8 bits)
S2: 62 (29.5 bits)
Wallclock time: 2 seconds
```

## ExPASy Home page Site Map Search ExPASy Contact us Proteomics tools Swiss-Prot

ESVEPEPEP	
sp   P50414   TGFB1_SHEEP	GQILSKLRLASPPSQGDVPPGPLPEAILALYNSTRDRVAG-ESAETEPEP
sp   019011   TGFB1_HORSE	GQILSKLRLASPPSQGEVPPGPLPEAVLALYNSTRAQVAG-ESAETEPEP
sp P18341 TGFB1 BOVIN	AILALYNSTRDRVAG-ESAETEPE <u>P</u> 🛰
sp P09531 TGFB1_CHICK	GQILSKLRLTAPPPASETPPRPLPDDVRALYNSTQELLKQRARLRPPPDG
· · · · · · · · · · · · · · · · · · ·	: *****:
sp P07200 TGFB1_PIG	EADYYAKEVTRVLMLESGNQIYDKFKGTPHSLYMLFNTSELREAVPEPVL

ExPASy Home page Site Map	Search ExPASy Contact us Proteomics tools Swiss-Prot
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SIB BLAST Netwo	rk Service William
This NCBI BLAST2 service is maintaine	ed by the Swiss Institute of Bioinformatics.
Click on the @ icons to access the online	BLAST help.
Accession number or sequence	
Enter a Swiss-Prot/TrEMBL accession n	number or a <b>PROTEIN</b> sequence in <b>RAW format</b> .
QHVELYQKYS NNS	
HTML 🔽	Output format:
Run BLAST or Reset Form	
Choose the appropriate BLAST	
• blastp - query against the UniProt	t Knowledgebase (Swiss-Prot + TrEMBL)
Taxonomic groups (not available for PD	B and translated EST):
select a database - Complete database - subsection	To restrict the search to a particular taxon, it is much faster to sele drop-down list on the left than to specify your own taxonomic group more accurate statistics.
or specify a taxonomic group	Enter a species name, a TaxID or the latin name of a taxonomic grows (";"). Example: Fungi; Homo sapiens.
	· · · · · · · · · · · · · · · · · · ·
or select a microbial proteome	

☐ Search only Swiss-Prot (curated sequences) ☐ Exclude fragment sequences
O blastp - query against another protein database UniRef100 Please, supply an email address if y
C tblastn - query against the six-frame translation of a nucleotide database All EMBL + GSS (without Taxonomic groups: All or select a microbial genome
② Your email address:  If an e-mail address is provided, results will be automatically mailed back (recommended for tblastn searches).  Run BLAST or Resel Form
Options:
<ul> <li>Comparison Matrix: Auto-select  </li> <li>Number of best scoring sequences to show: 100  </li> <li>Number of best alignments to show: 100  </li> <li>Setting the E threshold: 10  </li> <li>✓ (Number of expected matches in a random database)</li> <li>✓ Filter the sequence for low-complexity regions</li> <li>✓ Gapped alignment</li> <li>✓ Identity BLAST</li> </ul>
For a more 'advanced' BLAST2 search (with more options and programs, and a larger variety of databases) please use the <u>original submission form at EMBnet-CH (Lausanne)</u> .

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In case of problems, please read the <u>online BLAST help</u> .  If your question is not covered, please contact < <u>helpdesk@expasy.org</u> >.
NCBI BLAST program reference [PMID: 9254694]: Altschul S.F., Madden T.L., Schäffer A.A., Zhang J., Zhang Z., Miller W., Lipman D.J. Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. Nucleic Acids Res. 25:3389-3402(1997).
Query: 13 AA  Date run: 2005-09-24 15:53:30 UTC+0100 on sib-gml.unil.ch  Program: NCBI BLASTP 1.5.4-Paracel [2003-06-05]  Database: EXPASY/UniProtKB  2,312,053 sequences; 758,486,757 total letters  UniProt Knowledgebase Release 6.0 consists of:  UniProtKB/Swiss-Prot Release 48.0 of 13-Sep-2005: 194317 entries  UniProtKB/TrEMBL Release 31.0 of 13-Sep-2005: 2105517 entries
Taxonomic view Nice Blast view Printable view
List of potentially matching sequences
Send selected sequences to Clustal W (multiple alignment)
☐ Include query sequence
Db AC Description Score E-value
sp 000292 TGFB4 HUMAN Transforming growth factor beta 4 precurso 43 6e-04
tr <u>Q5TE89</u> _HUMAN Left-right determination factor 2 [LEFTY2] [Homo <u>43</u> 6e-04
tr <u>Q53H67</u> <u>HUMAN Left-right determination</u> , factor B preproprotein <u>43</u> 6e-04
sp 075610 LEFTB_HUMAN Left-right determination factor B precurso 38 0.019
tr Q5TE94 _HUMAN Left-right determination factor 1 [LEFTY1] [Homo 38 0.019
sp Q64280 TGFB4_MOUSE Transforming growth factor beta 4 precurso 38 0.025
sp P57785 LEFTB MOUSE Left-right determination factor B precurso 38 0.025
tr Q52M97 _XENLA Hypothetical protein [Xenopus laevis (African cl 38 0.025
tr Q8BMF7 MOUSE Mus musculus 13 days embryo male testis cDNA, RI 38 0.025

35 0.20 ☐ tr Q5UCE3 \_RAT EBAF precursor [Rattus norvegicus (Rat)] Graphical overview of the alignments to resubmit your query after masking regions matching PROSITE profiles Click here or Pfam HMMs (② Help) (use ScanProsite for more details about PROSITE matches) Profile hits Pfan hits **Hat** Matches on query sequence Submission 1 TGF84\_HUNAN Q5TE89 Q53M67 LEFTB\_HUNAN Q5TE94 TGF84\_HOUSE LEFTB\_HOUSE Q52M97 Q88MF7 Q5UCE3 Submission 75 100% 25 50 Identity Alignments Transforming growth factor beta 4 precursor (TGF-beta 4) 366 AA 000292 (Endometrial bleeding-associated factor) (Left-right TGFB4 HUMAN determination factor A) (Lefty-A protein) [EBAF] [Homo <u>align</u> sapiens (Human)] Score = 43.5 bits (95), Expect = 6e-04Identities = 13/13 (100%), Positives = 13/13 (100%) ASEASTHLLVFGM 13 Query: 1 **ASEASTHLLVFGM** Sbjct: 91 ASEASTHLLVFGM 103 Left-right determination factor 2 [LEFTY2] [Homo sapiens 366 tr Q5TE89

Query: 1 ASEASTHLLVFGM 13
ASEASTHLLVFGM

Sbjct: 91 ASEASTHLLVFGM 103

tr Q53H67 Left-right determination, factor B preproprotein variant Q53H67\_HUMAN (Fragment) [LEFTY1] [Homo sapiens (Human)] align Score = 43.5 bits (95), Expect = 6e-04Identities = 13/13 (100%), Positives = 13/13 (100%) Query: 1 ASEASTHLLVFGM 13 ASEASTHLLVFGM Sbjct: 91 ASEASTHLLVFGM 103 **sp** 075610 Left-right determination factor B precursor (Lefty-B 366 LEFTB\_HUMAN protein) AA [LEFTB] [Homo sapiens (Human)] align Score = 38.4 bits (83), Expect = 0.019Identities = 11/11 (100%), Positives = 11/11 (100%) Query: 3 EASTHLLVFGM 13 EASTHLLVFGM Sbjct: 93 EASTHLLVFGM 103 Left-right determination factor 1 [LEFTY1] [Homo sapiens 366 Q5TE94 HUMAN (Human)] AA align Score = 38.4 bits (83), Expect = 0.019Identities = 11/11 (100%), Positives = 11/11 (100%) EASTHLLVFGM 13 Query: 3 EASTHLLVFGM Sbjct: 93 EASTHLLVFGM 103 **sp** Q64280 Transforming growth factor beta 4 precursor (TGF-beta 4) 368 TGFB4\_MOUSE (Lefty AA protein) (Lefty-1 protein) (STRA3 protein) [Ebaf] [Mus align musculus (Mouse)] Score = 38.0 bits (82), Expect = 0.025Identities = 11/12 (91%), Positives = 11/12 (91%) SEASTHLLVFGM 13 Query: 2 SE STHLLVFGM Sbjct: 92 SETSTHLLVFGM 103

Left-right determination factor B precursor (Lefty-2 368 **sp** P57785 LEFTB\_MOUSE protein) AΑ [Leftb] [Mus musculus (Mouse)] <u>align</u> Score = 38.0 bits (82), Expect = 0.025Identities = 11/12 (91%), Positives = 11/12 (91%) SEASTHLLVFGM 13 Query: 2 SE STHLLVFGM Sbjct: 92 SETSTHLLVFGM 103 tr Q52M97 Hypothetical protein [Xenopus laevis (African clawed 368 Q52M97\_XENLA frog)] AA align Score = 38.0 bits (82), Expect = 0.025Identities = 11/12 (91%), Positives = 11/12 (91%) Query: 2 SEASTHLLVFGM 13 SE STHLLVFGM Sbjct: 92 SETSTHLLVFGM 103 tr Q8BMF7 Mus musculus 13 days embryo male testis cDNA, RIKEN 368 Q8BMF7 MOUSE full-length AΑ enriched library, clone: 6030463A22 product: LEFT-RIGHT align DETERMINATION FACTOR B (LEFTY-2 PROTEIN), full insert sequence (Left-right determination factor 2) [Lefty2] [Mus musculus (Mouse)] Score = 38.0 bits (82), Expect = 0.025Identities = 11/12 (91%), Positives = 11/12 (91%) Query: 2 SEASTHLLVFGM 13 SE STHLLVFGM Sbjct: 92 SETSTHLLVFGM 103 tr Q5UCE3 EBAF precursor [Rattus norvegicus (Rat)] 366 AA Q5UCE3 RAT align Score = 35.0 bits (75), Expect = 0.20Identities = 10/12 (83%), Positives = 10/12 (83%) Query: 2 SEASTHLLVFGM 13 SE S HLLVFGM Sbjct: 92 SETSSHLLVFGM 103

Database: EXPASY/UniProtKB

Posted date: Sep 12, 2005 12:34 PM

```
Number of letters in database: 758,486,757
 Number of sequences in database: 2,312,053
Lambda
           K
            0.277
   0.342
                      1.67
Gapped
Lambda
                     0.610
   0.294
            0.110
Matrix: PAM30
Gap Penalties: Existence: 9, Extension: 1
Number of HSP's successfully gapped in prelim test: 0
length of query: 13
length of database: 758,486,757
effective HSP length: 4
effective length of query: 9
effective length of database: 749,238,545
effective search space: 6743146905
effective search space used: 6743146905
T: 16
A: 40
X1: 15 ( 7.4 bits)
X2: 35 (14.8 bits)
X3: 58 (24.6 bits)
S1: 40 (21.6 bits)
S2: 62 (29.5 bits)
Wallclock time: 2 seconds
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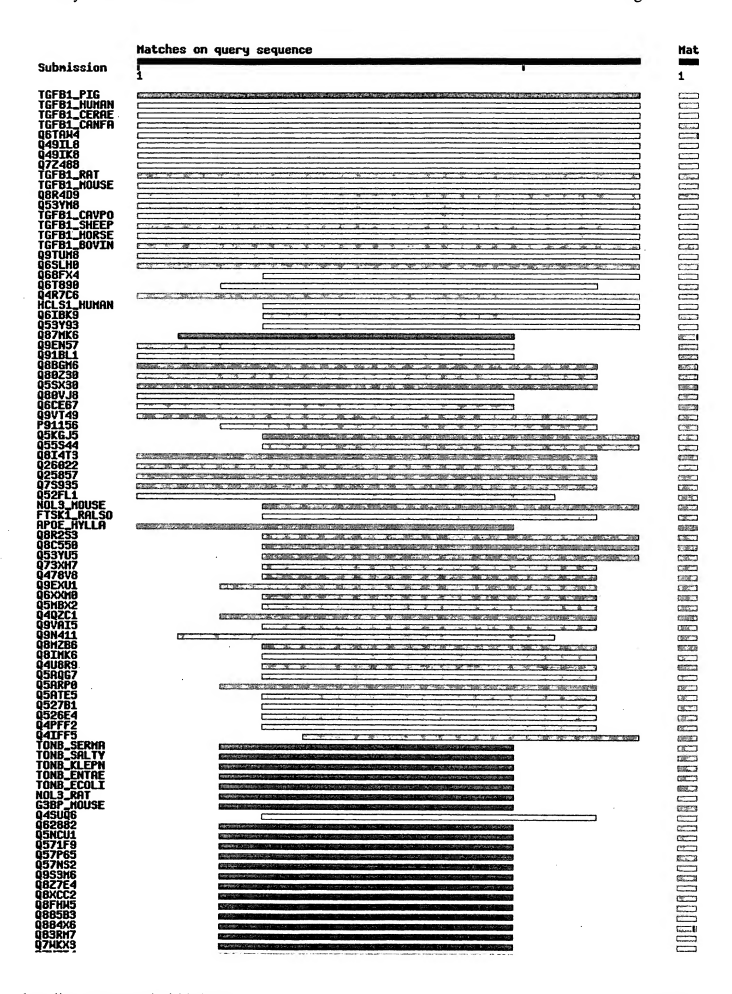
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the computation was performed at the SIB using the BLAST network service.  The SIB BLAST network service uses a server developed at SIB and the NCBI
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In case of problems, please read the <u>online BLAST help</u> .  If your question is not covered, please contact < <u>helpdesk@expasy.org</u> >.
NCBI BLAST program reference [PMID: 9254694]: Altschul S.F., Madden T.L., Schäffer A.A., Zhang J., Zhang Z., Miller W., Lipman D.J. Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. Nucleic Acids Res. 25:3389-3402(1997).
Query: 13 AA  Date run: 2005-09-24 15:46:19 UTC+0100 on sib-gm1.unil.ch  Program: NCBI BLASTP 1.5.4-Paracel [2003-06-05]  Database: EXPASY/UniProtKB  2,312,053 sequences; 758,486,757 total letters  UniProt Knowledgebase Release 6.0 consists of:  UniProtKB/Swiss-Prot Release 48.0 of 13-Sep-2005: 194317 entries  UniProtKB/TrEMBL Release 31.0 of 13-Sep-2005: 2105517 entries
Texamente view   NiceBlest view   Principle view
List of potentially matching sequences
Send selected sequences to Clustal W (multiple alignment)
☐ Include query sequence
Db AC Description Score E-value
sp P07200 TGFB1_PIG Transforming growth factor beta 1 precursor
☐ tr Q7Z488 _HUMAN Transforming growth factor beta 1 precursor (Fra 42 0.001 ☐ sp P17246 TGFB1_RAT Transforming growth factor beta 1 precursor 40 0.008

	cn P04202	TGFB1 MOUSE Transforming growth factor beta 1 precurso	40	0.008
		SIGHI Transforming growth factor beta-1 protein (Fragm		
		RAT TGF beta 1 [Rattus norvegicus (Rat)]		0.008
_		TGFB1 CAVPO Transforming growth factor beta 1 precurso		0.014
		<del>-</del>	—.	
		TGFB1_SHEEP Transforming growth factor beta 1 precurso		0.045
	_	TGFB1_HORSE Transforming growth factor beta 1 precurso		
		TGFB1_BOVIN Transforming growth factor beta 1 precurso		
		HORSE Transforming growth factor beta 1 [TGFb1] [Equus		
		_PERMA Transforming growth factor beta 1 (Fragment) [Tg		
		_RAT Hematopoietic cell specific Lyn substrate 1 (Predi		0.48
		_ACTPL TonB2 [tonB2] [Actinobacillus pleuropneumoniae (		0.64
		_MACFA Testis cDNA, clone: QtsA-15617, similar to human	<u>33</u>	0.64
	_	HCLS1_HUMAN Hematopoietic lineage cell specific protei	<u>33</u>	0.86
		_HUMAN HCLS1 protein [HCLS1] [Homo sapiens (Human)]	<u>33</u>	0.86
		_HUMAN Hematopoietic cell-specific Lyn substrate 1 [Hom		0.86
	tr <u>Q87MK6</u>	_VIBPA Hypothetical protein VP2226 [VP2226] [Vibrio par	<u>32</u>	1.1
	tr <u>Q9EN57</u>	_NPVST Zinc finger protein [Spodoptera litura multicaps	<u>32</u>	1.5
		_NPVST HOAR [Spodoptera litura multicapsid nucleopolyhe		1.5
	tr <u>Q8BGM6</u>	_MOUSE Mus musculus 10 days neonate cerebellum cDNA, RI	<u>32</u>	1.5
	tr <u>Q80Z30</u>	_RAT Calmodulin-dependent protein kinase phosphatase N	<u>32</u>	1.5
	tr <u>Q5SX30</u>	_MOUSE Protein phosphatase 1E (PP2C domain containing)	32	1.5
	tr <u>Q80VJ8</u>	_MOUSE Hypothetical protein BC050196 [BC050196] [Mus mu	<u>32</u>	2.1
	tr <u>Q6CE67</u>	_YARLI Similarity [YALIOB18172g] [Yarrowia lipolytica (	<u>32</u>	2.1
	tr <u>Q9VT49</u>	_DROME CG14168-PA [CG14168] [Drosophila melanogaster (F	<u>31</u>	2.8
		_CAEEL Hypothetical protein [C43E11.1] [Caenorhabditis		2.8
		_CRYNE Hypothetical protein [CNE03390] [Cryptococcus ne		2.8
		_CRYNE Hypothetical protein [CNBE3380] [Cryptococcus ne		2.8
		_PLAF7 PFG377 protein [PFL2405c] [Plasmodium falciparum		3.7
		_PLAFA Plasmodium falciparum sexual stage mRNA sequence	31	3.7
		_PLAFA Pfg377 [Plasmodium falciparum]	31	3.7
		_NEUCR Predicted protein [NCU07949.1] [Neurospora crassa]	<u>31</u>	3.7
		_MAGGR Hypothetical protein [MG01625.4] [Magnaporthe gr	<u>31</u>	3.7
		NOL3_MOUSE Nucleolar protein 3 [Nol3] [Mus musculus (M	30	5.0
	sp Q8XRHO	FTSK1_RALSO DNA translocase ftsK 1 [ftsK1] [Ralstonia	30	5.0
	sp Q9GLM6	APOE_HYLLA Apolipoprotein E precursor (Apo-E) [APOE] [	<u>30</u>	5.0
	•	_MOUSE Nucleolar protein 3 [Nol3] [Mus musculus (Mouse)]	<u>30</u>	5.0
	tr <u>Q8C550</u>	_MOUSE Mus musculus adult male hypothalamus cDNA, RIKEN	<u>30</u>	5.0
	tr <u>Q53YU5</u>	_MOUSE Apoptosis repressor interacting with CARD (0 day	<u>30</u>	5.0
	tr <u>Q73XH7</u>	_MYCPA Fas [fas] [Mycobacterium paratuberculosis]	<u>30</u>	5.0
	tr <u>Q478V8</u>	_9RHOO Response regulator receiver:CheW-like protein:AT	<u>30</u>	5.0
		_KLEPN TonB (Fragment) [tonB] [Klebsiella pneumoniae]	30	5.0
		_MYCSM Fatty acid synthetase I [fas1] [Mycobacterium sm	<u>30</u>	5.0
		_ACTSU TonB2 [tonB2] [Actinobacillus suis]	<u>30</u>	5.0
		_KLEPN Periplasmic energy transducer (Fragment) [tonB]	<u>30</u>	5.0
	tr <u>Q9VAI5</u>	_DROME CG15506-PB, isoform B [CG15506] [Drosophila mela	<u>30</u>	5.0

Г	tr Q9N411 _CAEEL Prion-like-(Q/n-rich)-domain-bearing protein pro	30	E 0
	<del></del>		5.0
			5.0
		_	5.0
	The state of the s		5.0
			5.0
			5.0
		<u>30</u>	5.0
		<u>30</u>	5.0
	tr Q526E4 _MAGGR Hypothetical protein [MG06862.4] [Magnaporthe gr	<u>30</u>	5.0
	<u> </u>		5.0
	tr Q4IFF5 _GIBZE Hypothetical protein [FG04053.1] [Gibberella zea	<u>30</u>	5.0
	sp P26185 TONB_SERMA TonB protein [tonB] [Serratia marcescens]	30	6.7
	sp P25945 TONB_SALTY TonB protein [tonB] [Salmonella typhimurium]	<u>30</u>	6.7
	sp P45610 TONB_KLEPN TonB protein [tonB] [Klebsiella pneumoniae]	<u>30</u>	6.7
	sp P46383 TONB_ENTAE TonB protein [tonB] [Enterobacter aerogenes	<u>30</u>	6.7
	sp P02929 TONB_ECOLI TonB protein [tonB] [Escherichia coli]	<u>30</u>	6.7
	sp Q62881 NOL3_RAT Nucleolar protein 3 [Nol3] [Rattus norvegicus	30	6.7
	sp P97855 G3BP_MOUSE Ras-GTPase-activating protein binding prote	30	6.7
	tr Q4SUQ6 _TETNG Chromosome undetermined SCAF13844, whole genome	30	6.7
	tr Q62882 _RAT Hypothetical protein [Rattus norvegicus (Rat)]	30	6.7
	tr Q5NCU1 MOUSE Ras-GTPase-activating protein SH3-domain binding	30	6.7
	tr Q571F9 MOUSE MKIAA4115 protein (Fragment) [G3bp] [Mus musculu	30	6.7
	tr Q57P65 SALCH SsrAB activated gene [srfA] [Salmonella cholerae	30	6.7
	tr <u>Q57NS2</u> <u>SALCH</u> Energy transducer; uptake of iron, cyanocobalimi	30	6.7
	tr Q9S3M6 BORPE TonB protein (Siderophore-mediated iron transpor	30	6.7
	tr Q8Z7E4 _SALTI TonB protein [tonB] [Salmonella typhi]	***************************************	6.7
	tr Q8XCC2 ECO57 Energy transducer; uptake of iron, cyanocobalimi	30	6.7
	tr Q8FHW5 _ECOL6 TonB protein [tonB] [Escherichia coli O6]	30	6.7
			6.7
		**********	6.7
			6.7
	tr Q7WKX3 _BORBR Siderophore-mediated iron transport protein [ton		6.7
	tr Q7W7I4 _BORPA Siderophore-mediated iron transport protein [ton		6.7
	tr Q5PHV4 _SALPA Putative virulence effector protein [srfA] [Salm	***********	6.7
	tr Q5PCS7 _SALPA TonB protein [tonB] [Salmonella paratyphi-a]	***************************************	6.7
	tr Q5NZS1 _AZOSE Translation initiation factor IF-2 [infB] [Azoar		6.7
	tr Q48GE4 _PSESH Flagellar assembly protein Flih [flih] [Pseudomo		
	tr Q9ZG18 BORBR TonB [tonB] [Bordetella bronchiseptica (Alcalige		6.7
	tr Q98329 _KLEPN TonB (Fragment) [tonB] [Klebsiella pneumoniae]		6.7
	tr Q98327 _KLEPN TonB (Fragment) [tonB] [Klebsiella pneumoniae]		6.7
			6.7
	· · · · · · · · · · · · · · · · · · ·	,	6.7
	<del> </del>		6.7
	· · · · · · · · · · · · · · · · · · ·		6.7
_	tr Q9R426 _KLEPN TonB (Fragment) [tonB] [Klebsiella pneumoniae]		6.7
<b></b> _	tr Q9EXU0 _KLEPN TonB (Fragment) [tonB] [Klebsiella pneumoniae]	<u>30</u>	6.7

tr Q9EXT9 _KLEPN TonB (Fragment) [tonB] [Klebsiella pneumoniae]	<u>30</u> 6.7
Graphical overview of the alignments	
click here to resubmit your query after masking regions matching PRO or Pfam HMMs	OSITE profiles
( Help) (use ScanProsite for more details about PROSITE	matches)
Profile hits	



#### Alignments

Transforming growth factor beta 1 precursor (TGF-beta 1) 390 **sp** P07200 TGFB1 PIG [TGFB1] AA [Sus scrofa (Pig)] align Score = 46.0 bits (101), Expect = 9e-05Identities = 13/13 (100%), Positives = 13/13 (100%) ESVEPEPEPEADY 13 Query: 1 ESVEPEPEPEADY Sbjct: 91 ESVEPEPEPEADY 103 **sp** P01137 Transforming growth factor beta 1 precursor (TGF-beta 1) 390 TGFB1\_HUMAN [TGFB1] AA[Homo sapiens (Human)] align Score = 42.2 bits (92), Expect = 0.001 Identities = 12/13 (92%), Positives = 12/13 (92%) **ESVEPEPEPEADY 13** Query: 1 ES EPEPEPEADY Sbjct: 91 ESAEPEPEPEADY 103 **sp** P09533 Transforming growth factor beta 1 precursor (TGF-beta 1) 390 TGFB1 CERAE [TGFB1] AA[Cercopithecus aethiops (Green monkey) (Grivet)] align Score = 42.2 bits (92), Expect = 0.001Identities = 12/13 (92%), Positives = 12/13 (92%) Query: 1 **ESVEPEPEPEADY 13** ES EPEPEPEADY Sbjct: 91 ESAEPEPEPEADY 103 **sp** P54831 Transforming growth factor beta 1 precursor (TGF-beta 1) 390 TGFB1 CANFA [TGFB1] AA [Canis familiaris (Dog)] align Score = 42.2 bits (92), Expect = 0.001Identities = 12/13 (92%), Positives = 12/13 (92%) Query: 1 ESVEPEPEPEADY 13 ES EPEPEPEADY Sbjct: 91 ESAEPEPEPEADY 103

tr Q6TAW4 TGF beta (Fragment) [Felis silvestris catus (Cat)] 226 AA Q6TAW4 FELCA align Score = 42.2 bits (92), Expect = 0.001Identities = 12/13 (92%), Positives = 12/13 (92%) Query: 1 ESVEPEPEPEADY 13 ES EPEPEPEADY Sbjct: 42 ESAEPEPEPEADY 54 tr Q49IL8 Transforming growth factor beta 1 precursor (Fragment) 118 Q49IL8 HUMAN [TGFB1] AΑ [Homo sapiens (Human)] align Score = 42.2 bits (92), Expect = 0.001 Identities = 12/13 (92%), Positives = 12/13 (92%) Query: 1 ESVEPEPEPEADY 13 ES EPEPEPEADY Sbjct: 91 ESAEPEPEPEADY 103 tr Q49IK8 Transforming growth factor beta 1 precursor (Fragment) 118 Q491K8 HUMAN [TGFB1] AΑ [Homo sapiens (Human)] align Score = 42.2 bits (92), Expect = 0.001 Identities = 12/13 (92%), Positives = 12/13 (92%) Query: 1 ESVEPEPEPEADY 13 ES EPEPEPEADY Sbjct: 91 ESAEPEPEPEADY 103 tr Q7Z488 Transforming growth factor beta 1 precursor (Fragment) 118 Q7Z488 HUMAN [TGFB1] AΑ [Homo sapiens (Human)] align Score = 42.2 bits (92), Expect = 0.001Identities = 12/13 (92%), Positives = 12/13 (92%) Query: 1 ESVEPEPEPEADY 13 ES EPEPEPEADY Sbjct: 91 ESAEPEPEPEADY 103 sp P17246 Transforming growth factor beta 1 precursor (TGF-beta 1) 390 TGFB1 RAT [Tgfb1] AA [Rattus norvegicus (Rat)] align

Score = 39.7 bits (86), Expect = 0.008Identities = 11/13 (84%), Positives = 12/13 (91%) ESVEPEPEPEADY 13 Query: 1 ES +PEPEPEADY Sbjct: 91 ESADPEPEPEADY 103 sp P04202 Transforming growth factor beta 1 precursor (TGF-beta 1) 390 TGFB1 MOUSE [Tgfb1] AΑ [Mus musculus (Mouse)] align Score = 39.7 bits (86), Expect = 0.008Identities = 11/13 (84%), Positives = 12/13 (91%) Query: 1 ESVEPEPEPEADY 13 ES +PEPEPEADY Sbjct: 91 ESADPEPEPEADY 103 tr. Q8R4D9 Transforming growth factor beta-1 protein (Fragment) 368 Q8R4D9 SIGHI [Tgfb1] AΑ [Sigmodon hispidus (Hispid cotton rat)] align Score = 39.7 bits (86), Expect = 0.008Identities = 11/13 (84%), Positives = 12/13 (91%) Query: 1 ESVEPEPEPEADY 13 ES +PEPEPEADY Sbjct: 69 ESADPEPEPEADY 81 tr Q53YM8 TGF beta 1 [Rattus norvegicus (Rat)] 390 AA Q53YM8\_RAT align Score = 39.7 bits (86), Expect = 0.008Identities = 11/13 (84%), Positives = 12/13 (91%) Query: 1 ESVEPEPEPEADY 13 ES +PEPEPEADY Sbjct: 91 ESADPEPEPEADY 103 Transforming growth factor beta 1 precursor (TGF-beta 1) 390 TGFB1\_CAVPO [TGFB1] AΑ [Cavia porcellus (Guinea pig)] align Score = 38.8 bits (84), Expect = 0.014Identities = 11/13 (84%), Positives = 11/13 (84%)

```
Query: 1
           ESVEPEPEPEADY 13
           ES EPEPEPE DY
Sbjct: 91 ESAEPEPEPDY 103
                Transforming growth factor beta 1 precursor (TGF-beta 1)
                                                                               390
   TGFB1_SHEEP [TGFB1]
                                                                               AΑ
                                                                               align
                 [Ovis aries (Sheep)]
 Score = 37.1 \text{ bits } (80), \text{ Expect} = 0.045
 Identities = 11/13 (84%), Positives = 11/13 (84%)
Query: 1
           ESVEPEPEPEADY 13
           ES E EPEPEADY
Sbjct: 91 ESAETEPEPEADY 103
sp 019011
                Transforming growth factor beta 1 precursor (TGF-beta 1)
                                                                               390
   TGFB1 HORSE [TGFB1]
                                                                               AA
                 [Equus caballus (Horse)]
                                                                               align
 Score = 37.1 bits (80), Expect = 0.045
 Identities = 11/13 (84%), Positives = 11/13 (84%)
           ESVEPEPEPEADY 13
Query: 1
           ES E EPEPEADY
Sbjct: 91 ESAETEPEPEADY 103
              Transforming growth factor beta 1 precursor (TGF-beta 1) 315 AA
   TGFB1 BOVIN (Fragment) [TGFB1] [Bos taurus (Bovine)]
                                                                              align
 Score = 37.1 \text{ bits } (80), \text{ Expect = } 0.045
 Identities = 11/13 (84%), Positives = 11/13 (84%)
Query: 1 ESVEPEPEPEADY 13
          ES E EPEPEADY
Sbjct: 16 ESAETEPEPEADY 28
tr 09TUM8
                 Transforming growth factor beta 1 [TGFb1] [Equus
                                                                               390
   Q9TUM8_HORSE caballus (Horse)]
                                                                               AA
                                                                               align
 Score = 37.1 \text{ bits } (80), \text{ Expect = } 0.045
 Identities = 11/13 (84%), Positives = 11/13 (84%)
          ESVEPEPEPEADY 13
Query: 1
           ES E EPEPEADY
Sbjct: 91 ESAETEPEPEADY 103
```

tr Q6SLH0 Transforming growth factor beta 1 (Fragment) [Tgfb1] 249 Q6SLH0\_PERMA [Peromyscus AA maniculatus (Deer mouse)] align Score = 36.7 bits (79), Expect = 0.061Identities = 10/13 (76%), Positives = 11/13 (83%) Query: 1 ESVEPEPEPEADY 13 ES +PEPEPE DY Sbjct: 57 ESADPEPEPETDY 69 tr Q68FX4 Hematopoietic cell specific Lyn substrate 1 (Predicted) 476 AA Q68FX4\_RAT [Hcls1\_predicted] [Rattus norvegicus (Rat)] <u>align</u> Score = 33.7 bits (72), Expect = 0.48Identities = 9/10 (90%), Positives = 9/10 (90%) Query: 4 EPEPEPEADY 13 EPEPEPE DY Sbjct: 369 EPEPEPEPDY 378 tr Q6T890 TonB2 [tonB2] [Actinobacillus pleuropneumoniae 285 Q6T890 ACTPL (Haemophilus AΑ pleuropneumoniae) ] align Score = 33.3 bits (71), Expect = 0.64Identities = 9/10 (90%), Positives = 10/10 (100%) Query: 3 VEPEPEPEAD 12 VEPEPEPEA+ Sbjct: 73 VEPEPEPEAE 82 tr Q4R7C6 Testis cDNA, clone: QtsA-15617, similar to human 475 Q4R7C6\_MACFA hematopoietic AΑ cell-specific Lyn substrate 1 (HCLS1),mRNA, RefSeq: align NM\_005335.3 [Macaca fascicularis (Crab eating macaque) (Cynomolgus monkey)] Score = 33.3 bits (71), Expect = 0.64Identities = 11/16 (68%), Positives = 11/16 (68%), Gaps = 3/16 (18%) Query: 1 ESV---EPEPEPEADY 13 E V EPEPEPE DY Sbjct: 357 EPVYEAEPEPEPENDY 372

**sp** P14317 Hematopoietic lineage cell specific protein 486 HCLS1 HUMAN (Hematopoietic AA cell-specific LYN substrate 1) (LckBP1) (p75) [HCLS1] <u>align</u> [Homo sapiens (Human)] Score = 32.9 bits (70), Expect = 0.86 Identities = 9/10 (90%), Positives = 9/10 (90%) Query: 4 EPEPEPEADY 13 EPEPEPE DY Sbjct: 369 EPEPEPENDY 378 tr Q6IBK9 HCLS1 protein [HCLS1] [Homo sapiens (Human)] 486 AA Q6IBK9 HUMAN align Score = 32.9 bits (70), Expect = 0.86Identities = 9/10 (90%), Positives = 9/10 (90%) Query: 4 EPEPEPEADY 13 EPEPEPE DY Sbjct: 369 EPEPEPENDY 378 tr Q53Y93 Hematopoietic cell-specific Lyn substrate 1 [Homo 486 Q53Y93 HUMAN sapiens (Human)] AA align Score = 32.9 bits (70), Expect = 0.86 Identities = 9/10 (90%), Positives = 9/10 (90%) Query: 4 EPEPEPEADY 13 EPEPEPE DY Sbjct: 369 EPEPEPENDY 378 Hypothetical protein VP2226 [VP2226] [Vibrio 370 Q87MK6\_VIBPA parahaemolyticus] AΑ align Score = 32.5 bits (69), Expect = 1.1Identities = 9/9 (100%), Positives = 9/9 (100%)

tr Q9EN57 Zinc finger protein [Spodoptera litura multicapsid 497 AA

Query: 2 SVEPEPEPE 10

Sbjct: 43 SVEPEPEPE 51

.SVEPEPEPE

```
Q9EN57_NPVST nucleopolyhedrovirus (SpltMNPV)]
                                                                     align
 Score = 32.0 bits (68), Expect = 1.5
 Identities = 9/10 (90%), Positives = 9/10 (90%)
Query: 1
          ESVEPEPEPE 10
          ES EPEPEPE
Sbjct: 355 ESAEPEPEPE 364
                 HOAR [Spodoptera litura multicapsid nucleopolyhedrovirus 731 AA
   Q91BL1_NPVST (SpltMNPV)]
                                                                           align
Score = 32.0 bits (68), Expect = 1.5
Identities = 9/10 (90%), Positives = 9/10 (90%)
          ESVEPEPEPE 10
Query: 1
           ES EPEPEPE
Sbjct: 459 ESAEPEPEPE 468
tr Q8BGM6
                Mus musculus 10 days neonate cerebellum cDNA, RIKEN
                                                                            151
   Q8BGM6 MOUSE full-length
                                                                            AΑ
                 enriched library, clone: B930008A12 product: similar to
                                                                            align
                 PP2CH (Mus musculus 0 day neonate eyeball cDNA, RIKEN
                 full-length enriched library, clone: E130208E03
                 product:similar to PP2CH) [Ppm1e] [Mus musculus
                 (Mouse) ]
 Score = 32.0 bits (68), Expect = 1.5
 Identities = 10/12 (83%), Positives = 11/12 (91%), Gaps = 1/12 (8%)
Query: 1 ESVEPEPEPEAD 12
          ES EPEPEPEA+
Sbjct: 36 ES-EPEPEPEAE 46
tr Q80Z30
               Calmodulin-dependent protein kinase phosphatase N [Ppmle]
                                                                            750
   Q80Z30 RAT [Rattus
                                                                            AA
               norvegicus (Rat)]
                                                                            align
Score = 32.0 bits (68), Expect = 1.5
Identities = 10/12 (83%), Positives = 11/12 (91%), Gaps = 1/12 (8%)
Query: 1 ESVEPEPEPEAD 12
          ES EPEPEPEA+
Sbjct: 36 ES-EPEPEPEAE 46
```

```
Protein phosphatase 1E (PP2C domain containing) [Ppm1e]
                                                                           749
tr Q5SX30
                                                                           AΑ
   Q5SX30 MOUSE [Mus
                                                                           align
                musculus (Mouse)]
 Score = 32.0 bits (68), Expect = 1.5
 Identities = 10/12 (83%), Positives = 11/12 (91%), Gaps = 1/12 (8%)
Query: 1 ESVEPEPEAD 12
         ES EPEPEPEA+
Sbjct: 36 ES-EPEPEPEAE 46
                Hypothetical protein BC050196 [BC050196] [Mus musculus
tr Q80VJ8
                                                                           578
   Q80VJ8 MOUSE (Mouse)]
                                                                           AΑ
                                                                           align
 Score = 31.6 bits (67), Expect = 2.1
 Identities = 9/10 (90%), Positives = 9/10 (90%)
Query: 1 ESVEPEPE 10
          E VEPEPEPE
Sbjct: 402 EEVEPEPEPE 411
                 Similarity [YALIOB18172g] [Yarrowia lipolytica (Candida 1080 AA
tr Q6CE67
   Q6CE67 YARLI lipolytica)]
                                                                          align
 Score = 31.6 bits (67), Expect = 2.1
 Identities = 9/10 (90%), Positives = 9/10 (90%)
          ESVEPEPEPE 10
Query: 1
           E VEPEPEPE
Sbjct: 116 EEVEPEPEPE 125
                                                                           734
tr 09VT49
                 CG14168-PA [CG14168] [Drosophila melanogaster (Fruit
                                                                           AA
   Q9VT49_DROME fly)]
                                                                           align
 Score = 31.2 bits (66), Expect = 2.8
 Identities = 10/12 (83%), Positives = 10/12 (83%)
           ESVEPEPEPEAD 12
Query: 1
           E VEPEPEPE D
Sbjct: 348 EVVEPEPEPEKD 359
                 Hypothetical protein [C43E11.1] [Caenorhabditis elegans] 580 AA
tr P91156
    P91156_CAEEL
                                                                           align
```

```
Score = 31.2 bits (66), Expect = 2.8
 Identities = 10/12 (83%), Positives = 10/12 (83%), Gaps = 2/12 (16%)
Query: 3
          VEPE--PEPEAD 12
          VEPE PEPEAD
Sbjct: 118 VEPEAEPEPEAD 129
                Hypothetical protein [CNE03390] [Cryptococcus neoformans 498 AA
tr Q5KGJ5
   Q5KGJ5 CRYNE (Filobasidiella neoformans)]
                                                                           align
 Score = 31.2 bits (66), Expect = 2.8
 Identities = 8/10 (80%), Positives = 9/10 (90%)
Query: 4 EPEPEPEADY 13
          EPEPEPE +Y
Sbjct: 300 EPEPEPEPEY 309
                Hypothetical protein [CNBE3380] [Cryptococcus neoformans 303
tr Q55S44
   Q55S44 CRYNE var.
                                                                           AA
                neoformans B-3501A]
                                                                           align
 Score = 31.2 bits (66), Expect = 2.8
 Identities = 8/10 (80%), Positives = 9/10 (90%)
Query: 4
         EPEPEPEADY 13
          EPEPEPE +Y
Sbjct: 111 EPEPEPEPY 120
tr Q8I4T3
                PFG377 protein [PFL2405c] [Plasmodium falciparum
                                                                          3119
   Q8I4T3_PLAF7 (isolate 3D7)]
                                                                          AA
                                                                          align
 Score = 30.8 bits (65), Expect = 3.7
 Identities = 10/13 (76%), Positives = 11/13 (83%), Gaps = 1/13 (7%)
Query: 1
          ESV-EPEPEPEAD 12
          E V EPEPEPEA+
Sbjct: 123 EEVREPEPEPEAE 135
tr Q26022
                Plasmodium falciparum sexual stage mRNA sequence.
                                                                           754
   Q26022 PLAFA (Fragment)
                                                                           AA
                 [Plasmodium falciparum]
                                                                           align
 Score = 30.8 bits (65), Expect = 3.7
 Identities = 10/13 (76%), Positives = 11/13 (83%), Gaps = 1/13 (7%)
```

Query: 1 ESV-EPEPEPEAD 12 E V EPEPEPEA+ Sbjct: 123 EEVREPEPEPEAE 135

Score = 30.8 bits (65), Expect = 3.7
Identities = 10/13 (76%), Positives = 11/13 (83%), Gaps = 1/13 (7%)

Query: 1 ESV-EPEPEPEAD 12 E V EPEPEPEA+ Sbjct: 123 EEVREPEPEPEAE 135

tr Q7S935 Predicted protein [NCU07949.1] [Neurospora crassa] 2826 AA Q7S935\_NEUCR

align

Score = 30.8 bits (65), Expect = 3.7 Identities = 10/13 (76%), Positives = 10/13 (76%), Gaps = 1/13 (7%)

Query: 1 ESV-EPEPEPED 12 E V EPEPEPE D Sbjct: 708 EPVAEPEPEPED 720

tr Q52FL1 Hypothetical protein [MG01625.4] [Magnaporthe grisea 6586 Q52FL1\_MAGGR 70-15] AA align

Score = 30.8 bits (65), Expect = 3.7 Identities = 10/11 (90%), Positives = 10/11 (90%), Gaps = 1/11 (9%)

Query: 1 ESVEPEPEA 11 ESV PEPEPEA

Sbjct: 4949 ESV-PEPEPEA 4958

sp Q9D1X0 Nucleolar protein 3 [No13] [Mus musculus (Mouse)] 220 AA NOL3\_MOUSE align

Score = 30.3 bits (64), Expect = 5.0
Identities = 8/10 (80%), Positives = 9/10 (90%)

Query: 4 EPEPEPEADY 13 EPEPEPE D+ Sbjct: 202 EPEPEPEPDF 211

```
DNA translocase ftsK 1 [ftsK1] [Ralstonia solanacearum
                                                                            959
   Q8XRH0
                                                                            AΑ
   FTSK1 RALSO (Pseudomonas
                                                                            align
                solanacearum)]
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/9 (88%), Positives = 9/9 (99%)
Query: 4
          EPEPEPEAD 12
           EPEPEPEA+
Sbjct: 334 EPEPEPEAE 342
               Apolipoprotein E precursor (Apo-E) [APOE] [Hylobates lar
                                                                            317
sp Q9GLM6
                                                                            AΑ
   APOE HYLLA (Common
                                                                            align
               qibbon)]
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/10 (80%), Positives = 9/10 (90%)
Query: 1 ESVEPEPEPE 10
          + VEPEPEPE
Sbjct: 22 QAVEPEPEPE 31
                 Nucleolar protein 3 [Nol3] [Mus musculus (Mouse)] 220 AA
tr Q8R2S3
   Q8R2S3 MOUSE
                                                                     align
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/10 (80%), Positives = 9/10 (90%)
Query: 4
           EPEPEPEADY 13
           EPEPEPE D+
Sbjct: 202 EPEPEPEPDF 211
                 Mus musculus adult male hypothalamus cDNA, RIKEN full-
                                                                            220
tr Q8C550
   Q8C550 MOUSE length
                                                                            AA
                 enriched library, clone:A230035L15 product:apoptosis
                                                                            align
                 repressor with CARD domain [No13] [Mus musculus (Mouse)]
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/10 (80%), Positives = 9/10 (90%)
Query: 4
           EPEPEPEADY 13
           EPEPEPE D+
Sbjct: 202 EPEPEPEPDF 211
```

Apoptosis repressor interacting with CARD (0 day neonate 220

tr Q53YU5

```
AA
   Q53YU5_MOUSE eyeball
                                                                            align
                 cDNA, RIKEN full-length enriched library,
                 clone:E130314L04 product:apoptosis repressor with CARD
                 domain) [Nol3] [Mus musculus (Mouse)]
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/10 (80%), Positives = 9/10 (90%)
Query: 4
          EPEPEPEADY 13
          EPEPEPE D+
Sbjct: 202 EPEPEPEPDF 211
tr Q73XH7
                Fas [fas] [Mycobacterium paratuberculosis] 3092 AA
   Q73XH7_MYCPA
                                                             align
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/9 (88%), Positives = 9/9 (99%)
            EPEPEPEAD 12
Query: 4
            +PEPEPEAD
Sbjct: 1744 DPEPEPEAD 1752
                 Response regulator receiver: CheW-like protein: ATP-
                                                                           1866
tr Q478V8
                                                                           AA
   Q478V8 9RHOO binding region,
                 ATPase-like: Hpt [Daro 3895] [Dechloromonas aromatica
                                                                           align
                 RCB]
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/9 (88%), Positives = 9/9 (99%)
Query: 4
          EPEPEPEAD 12
           EPEPEPEA+
Sbjct: 831 EPEPEPEAE 839
                 TonB (Fragment) [tonB] [Klebsiella pneumoniae] 180 AA
tr Q9EXU1
   Q9EXU1 KLEPN
                                                                  align
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/10 (80%), Positives = 9/10 (90%)
Query: 3 VEPEPEPEAD 12
          VEPEPEPE +
Sbjct: 42 VEPEPEPETE 51
```

Fatty acid synthetase I [fas1] [Mycobacterium

tr Q6XXM0

3089

```
AA
   Q6XXM0_MYCSM smegmatis]
                                                                            align
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/9 (88%), Positives = 9/9 (99%)
Query: 4
            EPEPEPEAD 12
            +PEPEPEAD
Sbjct: 1747 DPEPEPEAD 1755
tr Q5MBX2
                 TonB2 [tonB2] [Actinobacillus suis] 281 AA
   Q5MBX2 ACTSU
                                                       align
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/9 (88%), Positives = 9/9 (99%)
Query: 4 EPEPEPEAD 12
          EPEPEPEA+
Sbjct: 74 EPEPEPEAE 82
tr Q4QZC1
                 Periplasmic energy transducer (Fragment) [tonB]
                                                                             138
   Q4QZC1_KLEPN [Klebsiella
                                                                             AΑ
                 pneumoniae]
                                                                             align
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/10 (80%), Positives = 9/10 (90%)
Query: 3 VEPEPEPEAD 12
          VEPEPEPE +
Sbjct: 21 VEPEPEPETE 30
tr Q9VAI5
                 CG15506-PB, isoform B [CG15506] [Drosophila melanogaster
                                                                             336
   Q9VAI5 DROME (Fruit
                                                                             AA
                 fly)]
                                                                             <u>align</u>
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/9 (88%), Positives = 9/9 (99%)
Query: 4
           EPEPEPEAD 12
           EPEPEPEA+
Sbjct: 321 EPEPEPEAE 329
tr Q9N411
                 Prion-like-(Q/n-rich)-domain-bearing protein protein 82
                                                                             300
   Q9N411_CAEEL [pqn-82]
                                                                             AΑ
                 [Caenorhabditis elegans]
                                                                             align
 Score = 30.3 bits (64), Expect = 5.0
```

```
Identities = 9/10 (90%), Positives = 9/10 (90%)
Query: 2
           SVEPEPEPA 11
           SVE EPEPEA
Sbjct: 244 SVEQEPEPEA 253
                 AT15667p [CG15506] [Drosophila melanogaster (Fruit fly)] 297 AA
tr Q8MZB6
   Q8MZB6_DROME
                                                                           align
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/9 (88%), Positives = 9/9 (99%)
Query: 4
          EPEPEPEAD 12
           EPEPEPEA+
Sbjct: 282 EPEPEPEAE 290
                 CG15506-PA, isoform A [CG15506] [Drosophila melanogaster 297
tr Q8IMK6
   Q8IMK6 DROME (Fruit
                                                                            AΑ
                                                                            align
                 fly)]
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/9 (88%), Positives = 9/9 (99%)
          EPEPEPEAD 12
Query: 4
           EPEPEPEA+
Sbjct: 282 EPEPEPEAE 290
tr Q4U8R9
                 Hypothetical protein [TA10045] [Theileria annulata] 599 AA
   Q4U8R9_THEAN
                                                                      align
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/9 (88%), Positives = 9/9 (99%)
Query: 4 EPEPEPEAD 12
          EPEPEPEA+
Sbjct: 315 EPEPEPEAE 323
tr Q5AQG7
                 Hypothetical protein [AN9463.2] [Aspergillus nidulans
                                                                           1737
   Q5AQG7 EMENI FGSC A4]
                                                                           AΑ
                                                                           align
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/9 (88%), Positives = 9/9 (99%)
Query: 4
          EPEPEPEAD 12
```

```
EPEPEPEA+
Sbjct: 448 EPEPEPEAE 456
```

```
Predicted protein [AN9040.2] [Aspergillus nidulans FGSC
tr Q5ARP0
                                                                          179
   Q5ARPO EMENI A4]
                                                                          AΑ
                                                                          align
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 10/14 (71%), Positives = 10/14 (71%), Gaps = 4/14 (28%)
Query: 3 VEPEPE----PEAD 12
          VEPEPE PEAD
Sbjct: 95 VEPEPEPEPAPEAD 108
tr Q5ATE5
                Hypothetical protein [AN8435.2] [Aspergillus nidulans
                                                                          850
   Q5ATE5 EMENI FGSC A4]
                                                                          AΑ
                                                                          align
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/9 (88%), Positives = 9/9 (99%)
Query: 4 EPEPEPEAD 12
          EPEPEPEA+
Sbjct: 661 EPEPEPEAE 669
                Hypothetical protein [MG06645.4] [Magnaporthe grisea 70- 717
tr Q527B1
   Q527B1 MAGGR 15]
                                                                          AA
                                                                          align
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/9 (88%), Positives = 9/9 (99%)
Query: 4 EPEPEPEAD 12
          EPEPEPEA+
Sbjct: 356 EPEPEPEAE 364
tr Q526E4
                Hypothetical protein [MG06862.4] [Magnaporthe grisea 70- 919
   Q526E4 MAGGR 15]
                                                                          AA
                                                                          align
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/9 (88%), Positives = 9/9 (99%)
Query: 4 EPEPEPEAD 12
          EPEPEPEA+
Sbjct: 466 EPEPEPEAE 474
```

```
tr <u>Q4P</u>FF2
                 Hypothetical protein [UM01161.1] [Ustilago maydis 521] 2649 AA
   Q4PFF2_USTMA
                                                                            align
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/9 (88%), Positives = 9/9 (99%)
Query: 4
            EPEPEPEAD 12
            EPEPEPEA+
Sbjct: 2312 EPEPEPEAE 2320
                 Hypothetical protein [FG04053.1] [Gibberella zeae PH-1] 807 AA
tr Q4IFF5
   Q4IFF5 GIBZE
                                                                             align
 Score = 30.3 bits (64), Expect = 5.0
 Identities = 8/9 (88%), Positives = 8/9 (88%)
           PEPEPEADY 13
Query: 5
           PEPEPE DY
Sbjct: 236 PEPEPEPDY 244
               TonB protein [tonB] [Serratia marcescens] 247 AA
sp P26185
   TONB_SERMA
                                                            <u>align</u>
 Score = 29.9 bits (63), Expect = 6.7
 Identities = 8/8 (100%), Positives = 8/8 (100%)
Query: 3 VEPEPEPE 10
          VEPEPEPE
Sbjct: 75 VEPEPEPE 82
sp P25945
               TonB protein [tonB] [Salmonella typhimurium] 242 AA
   TONB_SALTY
                                                               align
 Score = 29.9 \text{ bits } (63), \text{ Expect = } 6.7
 Identities = 8/8 (100%), Positives = 8/8 (100%)
Query: 3 VEPEPEPE 10
          VEPEPEPE
Sbjct: 69 VEPEPEPE 76
```

TonB protein [tonB] [Klebsiella pneumoniae] 243 AA

sp P45610

TONB\_KLEPN

```
align
```

Score = 29.9 bits (63), Expect = 6.7 Identities = 8/8 (100%), Positives = 8/8 (100%) Query: 3 VEPEPEPE 10 VEPEPEPE Sbjct: 68 VEPEPEPE 75 TonB protein [tonB] [Enterobacter aerogenes (Aerobacter 243 AA **sp** P46383 TONB ENTAE aerogenes)] align Score = 29.9 bits (63), Expect = 6.7Identities = 8/8 (100%), Positives = 8/8 (100%) Query: 3 VEPEPEPE 10 VEPEPEPE Sbjct: 69 VEPEPEPE 76 TonB protein [tonB] [Escherichia coli] 239 AA **sp** P02929 TONB ECOLI align Score = 29.9 bits (63), Expect = 6.7Identities = 8/8 (100%), Positives = 8/8 (100%) Query: 3 VEPEPEPE 10 VEPEPEPE Sbjct: 69 VEPEPEPE 76 **sp** Q62881 Nucleolar protein 3 [Nol3] [Rattus norvegicus (Rat)] 221 AA NOL3 RAT align Score = 29.9 bits (63), Expect = 6.7Identities = 8/8 (100%), Positives = 8/8 (100%) VEPEPEPE 10 Query: 3 VEPEPEPE Sbjct: 179 VEPEPEPE 186 **sp** P97855 Ras-GTPase-activating protein binding protein 1 (GAP SH3-465 G3BP MOUSE domain AΑ binding protein 1) (G3BP-1) (DNA helicase VIII) <u>align</u> (HDH-VIII) [G3bp] [Mus musculus (Mouse)]

Score = 29.9 bits (63), Expect = 6.7

```
Identities = 8/8 (100%), Positives = 8/8 (100%)
Query: 3
           VEPEPEPE 10
           VEPEPEPE
Sbjct: 191 VEPEPEPE 198
                  Chromosome undetermined SCAF13844, whole genome shotgun
                                                                                 129
tr Q4SUQ6
   Q4SUQ6 TETNG sequence
                                                                                 AΑ
                  [GSTENG00012352001] [Tetraodon nigroviridis (Green
                                                                                 align
                  puffer)]
 Score = 29.9 \text{ bits } (63), \text{ Expect = } 6.7
 Identities = 8/9 (88%), Positives = 8/9 (88%)
Query: 4 EPEPEPEAD 12
          EPEPEPE D
Sbjct: 90 EPEPEPETD 98
tr Q62882
                Hypothetical protein [Rattus norvegicus (Rat)] 139 AA
   Q62882_RAT
                                                                   align
 Score = 29.9 \text{ bits } (63), \text{ Expect = } 6.7
 Identities = 8/8 (100%), Positives = 8/8 (100%)
           VEPEPEPE 10
Query: 3
           VEPEPEPE
Sbjct: 97 VEPEPEPE 104
tr Q5NCU1
                  Ras-GTPase-activating protein SH3-domain binding protein 465 AA
   Q5NCU1_MOUSE [RP23-336J1.4] [Mus musculus (Mouse)]
                                                                                 align
 Score = 29.9 \text{ bits } (63), \text{ Expect = } 6.7
 Identities = 8/8 (100%), Positives = 8/8 (100%)
           VEPEPEPE 10
Query: 3
           VEPEPEPE
Sbjct: 191 VEPEPEPE 198
tr Q571F9
                  MKIAA4115 protein (Fragment) [G3bp] [Mus musculus
                                                                                 505
   Q571F9 MOUSE (Mouse)]
                                                                                 AA
                                                                                 align
 Score = 29.9 \text{ bits } (63), \text{ Expect = } 6.7
 Identities = 8/8 (100%), Positives = 8/8 (100%)
```

Query: 3 VEPEPEPE 10 VEPEPEPE

Sbjct: 231 VEPEPEPE 238

tr  $\frac{\text{Q57P65}}{\text{Q57P65}}$  SsrAB activated gene [srfA] [Salmonella choleraesuis] 441 AA  $\frac{\text{Q57P65}}{\text{Q57P65}}$  SALCH

align

Score = 29.9 bits (63), Expect = 6.7
Identities = 8/8 (100%), Positives = 8/8 (100%)

Query: 3 VEPEPEPE 10 VEPEPEPE

Sbjct: 169 VEPEPEPE 176

tr Q57NS2 Energy transducer; uptake of iron, cyanocobalimin; 281
Q57NS2\_SALCH sensitivity to AA
phages, colicins [tonB] [Salmonella choleraesuis] align

Score = 29.9 bits (63), Expect = 6.7 Identities = 8/8 (100%), Positives = 8/8 (100%)

Query: 3 VEPEPEPE 10 VEPEPEPE

Sbjct: 110 VEPEPEPE 117

tr Q9S3M6 TonB protein (Siderophore-mediated iron transport 266 Q9S3M6\_BORPE protein) [tonB] AA [Bordetella pertussis] align

Score = 29.9 bits (63), Expect = 6.7
Identities = 8/8 (100%), Positives = 8/8 (100%)

Query: 3 VEPEPEPE 10 VEPEPEPE

Sbjct: 101 VEPEPEPE 108

tr <u>Q8Z7E4</u> TonB protein [tonB] [Salmonella typhi] 242 AA Q8Z7E4\_SALTI align

Score = 29.9 bits (63), Expect = 6.7
Identities = 8/8 (100%), Positives = 8/8 (100%)

Query: 3 VEPEPEPE 10 VEPEPEPE

Sbjct: 69 VEPEPEPE 76

tr Q8XCC2 Energy transducer; uptake of iron, cyanocobalimin; 239
Q8XCC2\_EC057 sensitivity to AA
phages, colicins [tonB] [Escherichia coli 0157:H7] align

Score = 29.9 bits (63), Expect = 6.7
Identities = 8/8 (100%), Positives = 8/8 (100%)

Query: 3 VEPEPEPE 10 VEPEPEPE 56 VEPEPEPE 76

tr Q8FHW5 TonB protein [tonB] [Escherichia coli O6] 255 AA Q8FHW5\_ECOL6

align

Score = 29.9 bits (63), Expect = 6.7
Identities = 8/8 (100%), Positives = 8/8 (100%)

Query: 3 VEPEPEPE 10 VEPEPEPE Sbjct: 86 VEPEPEPE 93

tr Q885B3 Major facilitator family transporter [PSPTO1922] 465
Q885B3\_PSESM [Pseudomonas syringae pv. tomato] align

Score = 29.9 bits (63), Expect = 6.7
Identities = 8/8 (100%), Positives = 8/8 (100%)

Query: 3 VEPEPEPE 10 VEPEPEPE Sbjct: 422 VEPEPEPE 429

tr <u>Q884X6</u> Flagellar assembly protein Flih, putative [PSPT01960] 272 AA Q884X6\_PSESM [Pseudomonas syringae pv. tomato]

<u>align</u>

Score = 29.9 bits (63), Expect = 6.7 Identities = 8/8 (100%), Positives = 8/8 (100%)

Query: 3 VEPEPEPE 10 VEPEPEPE 50 VEPEPEPE 40

242 Membrane protein, energy transducer [tonB] [Shigella tr Q83RM7 AΑ Q83RM7\_SHIFL flexneri] align Score = 29.9 bits (63), Expect = 6.7Identities = 8/8 (100%), Positives = 8/8 (100%) Query: 3 VEPEPEPE 10 VEPEPEPE Sbjct: 72 VEPEPEPE 79 tr Q7WKX3 Siderophore-mediated iron transport protein [tonB] 268 Q7WKX3 BORBR [Bordetella AΑ bronchiseptica (Alcaligenes bronchisepticus)] align Score = 29.9 bits (63), Expect = 6.7Identities = 8/8 (100%), Positives = 8/8 (100%) Query: 3 VEPEPEPE 10 VEPEPEPE Sbjct: 103 VEPEPEPE 110 278 Siderophore-mediated iron transport protein [tonB] Q7W7I4 BORPA [Bordetella AΑ parapertussis] align Score = 29.9 bits (63), Expect = 6.7Identities = 8/8 (100%), Positives = 8/8 (100%) Query: 3 VEPEPEPE 10 VEPEPEPE Sbjct: 113 VEPEPEPE 120 Score = 29.9 bits (63), Expect = 6.7Identities = 8/8 (100%), Positives = 8/8 (100%) Query: 3 VEPEPEPE 10 VEPEPEPE Sbjct: 103 VEPEPEPE 110 tr Q5PHV4 Putative virulence effector protein [srfA] [Salmonella 399 AA Q5PHV4\_SALPA paratyphi-a] align Score = 29.9 bits (63), Expect = 6.7Identities = 8/8 (100%), Positives = 8/8 (100%) VEPEPEPE 10 Query: 3 VEPEPEPE

Sbjct: 127 VEPEPEPE 134

tr <u>Q5PCS7</u> TonB protein [tonB] [Salmonella paratyphi-a] 242 AA Q5PCS7\_SALPA

align

Score = 29.9 bits (63), Expect = 6.7 Identities = 8/8 (100%), Positives = 8/8 (100%)

Query: 3 VEPEPEPE 10 VEPEPEPE 56 VEPEPEPE 76

tr Q5NZS1 Translation initiation factor IF-2 [infB] [Azoarcus sp. 945 Q5NZS1\_AZOSE (strain EbN1)] AA

Score = 29.9 bits (63), Expect = 6.7
Identities = 8/8 (100%), Positives = 8/8 (100%)

Query: 3 VEPEPEPE 10 VEPEPEPE Sbjct: 159 VEPEPEPE 166

tr Q48GE4 Flagellar assembly protein Flih [fliH] [Pseudomonas 274 Q48GE4\_PSESH syringae pv. AA phaseolicola 1448A] align

Score = 29.9 bits (63), Expect = 6.7
Identities = 8/8 (100%), Positives = 8/8 (100%)

Query: 3 VEPEPEPE 10 VEPEPEPE Sbjct: 33 VEPEPEPE 40

tr Q9ZG18 TonB [tonB] [Bordetella bronchiseptica (Alcaligenes 268 AA Q9ZG18\_BORBR bronchisepticus)]

align

Score = 29.9 bits (63), Expect = 6.7 Identities = 8/8 (100%), Positives = 8/8 (100%)

Query: 3 VEPEPEPE 10 VEPEPEPE Sbjct: 103 VEPEPEPE 110

```
TonB (Fragment) [tonB] [Klebsiella pneumoniae] 180 AA
tr <u>Q9S3Z9</u>
   Q9S3Z9 KLEPN
                                                                    align
 Score = 29.9 bits (63), Expect = 6.7
 Identities = 8/8 (100%), Positives = 8/8 (100%)
Query: 3 VEPEPEPE 10
          VEPEPEPE
Sbjct: 42 VEPEPEPE 49
                 TonB (Fragment) [tonB] [Klebsiella pneumoniae] 180 AA
tr Q9S3Z7
   Q9S3Z7_KLEPN
                                                                    align
 Score = 29.9 \text{ bits } (63), \text{ Expect = } 6.7
 Identities = 8/8 (100%), Positives = 8/8 (100%)
Query: 3 VEPEPEPE 10
          VEPEPEPE
Sbjct: 42 VEPEPEPE 49
tr Q9S3Z6
                 TonB (Fragment) [tonB] [Klebsiella pneumoniae] 180 AA
   Q9S3Z6_KLEPN
                                                                    align
 Score = 29.9 \text{ bits } (63), \text{ Expect = } 6.7
 Identities = 8/8 (100%), Positives = 8/8 (100%)
Query: 3 VEPEPEPE 10
          VEPEPEPE
Sbjct: 42 VEPEPEPE 49
tr Q9R440
                  TonB (Fragment) [tonB] [Klebsiella pneumoniae] 180 AA
   Q9R440 KLEPN
                                                                    align
 Score = 29.9 bits (63), Expect = 6.7
 Identities = 8/8 (100%), Positives = 8/8 (100%)
Query: 3 VEPEPEPE 10
          VEPEPEPE
Sbjct: 42 VEPEPEPE 49
tr Q9R427
                 TonB (Fragment) [tonB] [Klebsiella pneumoniae] 180 AA
   Q9R427_KLEPN
                                                                    align
```

```
Score = 29.9 \text{ bits } (63), \text{ Expect = } 6.7
   Identities = 8/8 (100%), Positives = 8/8 (100%)
  Query: 3 VEPEPEPE 10
            VEPEPEPE
  Sbjct: 42 VEPEPEPE 49
                    TonB (Fragment) [tonB] [Klebsiella pneumoniae] 180 AA
 tr Q9R426
     Q9R426 KLEPN
                                                                       align
  Score = 29.9 \text{ bits } (63), \text{ Expect = } 6.7
   Identities = 8/8 (100%), Positives = 8/8 (100%)
 Query: 3 VEPEPEPE 10
            VEPEPEPE
 Sbjct: 42 VEPEPEPE 49
 tr Q9EXU0
                    TonB (Fragment) [tonB] [Klebsiella pneumoniae] 180 AA
     Q9EXU0 KLEPN
                                                                       align
  Score = 29.9 \text{ bits } (63), \text{ Expect = } 6.7
  Identities = 8/8 (100%), Positives = 8/8 (100%)
 Query: 3 VEPEPEPE 10
            VEPEPEPE
  Sbjct: 42 VEPEPEPE 49
 tr <u>Q9EXT</u>9
                   TonB (Fragment) [tonB] [Klebsiella pneumoniae] 180 AA
     Q9EXT9 KLEPN
                                                                       align
  Score = 29.9 \text{ bits } (63), \text{ Expect = } 6.7
  Identities = 8/8 (100%), Positives = 8/8 (100%)
 Query: 3 VEPEPEPE 10
            VEPEPEPE
 Sbjct: 42 VEPEPEPE 49
Database: EXPASY/UniProtKB
    Posted date: Sep 12, 2005 12:34 PM
  Number of letters in database: 758,486,757
  Number of sequences in database: 2,312,053
Lambda
          K
   0.319
         0.283
                      1.59
Gapped
```

```
Lambda
           K
   0.294
           0.110
                     0.610
Matrix: PAM30
Gap Penalties: Existence: 9, Extension: 1
length of query: 13
length of database: 758,486,757
effective HSP length: 4
effective length of query: 9
effective length of database: 749,238,545
effective search space: 6743146905
effective search space used: 6743146905
T: 16
A: 40
X1: 16 (7.4 bits)
X2: 35 (14.8 bits)
X3: 58 (24.6 bits)
S1: 43 (21.6 bits)
S2: 62 (29.5 bits)
Wallclock time: 3 seconds
```

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**Expanded** Expanding Expansion Expan

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Clear

# UniProtKB/TrEMBL entry Q53H67

Printer-trianally view

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[Entry info] [Name and origin] [References] [Comments] [Cross-references] [Keywords] [Features] [Sequence] [Tools]

Note: most headings are clickable, even if they don't appear as links. They link to the user manual or other documents.

**Entry information** 

Entry name

Q53H67 HUMAN

Primary accession number

**Q53H67** 

Secondary accession numbers

None

Entered in TrEMBL in

Release 31, September 2005

Sequence was last modified in

Release 31, September 2005.

Annotations were last modified in

Release 31, September 2005

Name and origin of the protein

Protein name

Left-right determination, factor B preproprotein variant

[Fragment]

Synonyms

None

Gene name

Name: LEFTY1

From

Homo sapiens (Human) [TaxID: 9606]

**Taxonomy** 

Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;

Mammalia; Eutheria; Euarchontoglires; Primates; Catarrhini;

Hominidae; Homo.

## References

# [1] NUCLEOTIDE SEQUENCE.

TISSUE=Colon:

Maruyama K., Sugano S.;

"Oligo-capping: a simple method to replace the cap structure of eucaryotic mRNAs with oligoribonucleotides.":

Gene 138:171-174(1994).

## [2] NUCLEOTIDE SEQUENCE.

TISSUE=Colon;

Suzuki Y., Yoshitomo K., Maruyama K., Suyama A., Sugano S.;

"Construction and characterization of a full length-enriched and a 5'-end-enriched cDNA library."; Gene 200:149-156(1997).

[3] NUCLEOTIDE SEOUENCE.

TISSUE=Colon:

Suzuki Y., Sugano S., Totoki Y., Toyoda A., Takeda T., Sakaki Y., Tanaka A., Yokoyama S.; Submitted (APR-2005) to the EMBL/GenBank/DDBJ databases.

#### Comments

• **SIMILARITY**: Belongs to the TGF-beta family.

#### **Cross-references**

**EMBL** 

AK222714; BAD96434.1; -; mRNA.[EMBL / GenBank / DDBJ] [CoDingSequence]

Ensembl ENSG00000143787; Homo sapiens. [Contig view]

HGNC HGNC:6552; LEFTY1. CleanEx HGNC:6552; LEFTY1.

GeneCards LEFTY1.

GeneLynx LEFTY1; Homo sapiens.

GenAtlas LEFTY1.

GO

GO:0008083; Molecular function: growth factor activity (inferred from electronic

annotation).

GO:0005160; Molecular function: transforming growth factor beta receptor binding

(inferred from electronic annotation).

GO:0016049; Biological process: cell growth (inferred from electronic annotation).

QuickGo view.

IPR001839; TGFb. IPR003942; TGFb4.

InterPro IPR001111; TGFb N.

Graphical view of domain structure.

PF00019; TGF beta; 1.

Pfam PF00688; TGFb\_propeptide; 1.

Pfam graphical view of domain structure.

PRINTS PR01427; TGFBETA4.

ProDom PD000357; TGFb; 1.

[Domain structure / List of seq. sharing at least 1 domain]

SMART SM00204; TGFB; 1.

PROSITE PS00250; TGF\_BETA\_1; 1. HOVERGEN [Family / Alignment / Tree]

ProtoMap Q53H67. PRESAGE Q53H67. ModBase Q53H67.

SWISS-2DPAGE

Get region on 2D PAGE.

UniRef View cluster of proteins with at least 50% / 90% identity.

# Keywords

#### Growth factor.

# Features



Feature table viewer

Key From To Length Description

NON\_TER 1 1

#### Sequence information

Length: 366 AA [This is the length of the partial sequence]

Molecular weight: 40826 Da [This is the MW of the partial sequence]

CRC64: A5AA26DE37658075 [This is a checksum on the sequence]

 $1\underline{0}$   $2\underline{0}$   $3\underline{0}$   $4\underline{0}$   $5\underline{0}$   $6\underline{0}$ 

MQPLWLCWAL WVLPLASPGA ALTGEQLLGS LLRQLQLKEV PTLDRADMEE LVIPTHVRAQ 90 100 YVALLQRSHG DRSRGKRFSQ SFREVAGRFL ASEASTHLLV FGMEQRLPPN SELVQAVLRL FQEPVPKAAL HRHGRLSPRS AQARVTVEWL RVRDDGSNRT SLIDSRLVSV HESGWKAFDV 200 210 230 -TEAVNFWQQL SRPRQPLLLQ VSVQREHLGP LASGAHKLVR FASQGAPAGL GEPQLELHTL 270 280 DLGDYGAQGD CDPEAPMTEG TRCCRQEMYI DLQGMKWAEN WVLEPPGFLA YECVGTCRQP 310 320 330 340 350 360 PEALAFKWPF LGPRQCIASE TDSLPMIVSI KEGGRTRPQV VSLPNMRVQK CSCASDGALV

PRRLQP

Q53H67 in FASTA format

View entry in original UniProtKB/TrEMBL format View entry in raw text format (no links) Request for annotation of this UniProtKB/TrEMBL entry

**BLAST** ExPASy/SIB

BLAST submission on or at NCBI (USA)



Sequence analysis tools: ProtParam, ProtScale, Compute pI/Mw, PeptideMass, PeptideCutter, Dotlet (Java)



ScanProsite, MotifScan



Submit a homology modeling request to SWISS-MODEL



NPSA Sequence analysis tools

**ExPASy Home page** Site Map Search ExPASy Contact us **Swiss-Prot** Hosted by NCSC US Mirror sites: Australia Brazil Canada Korea Switzerland Taiwan

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**UMBER Home** 

Contents

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Advanced Search

Relation Search

# ==SPRINT==> PRINTS View



#### PR01427

Identifier

TGFBETA4 [View Relations] [View Alignment]

Accession

PR01427

No. of Motifs

Creation Date

01-NOV-2000

Title

Transforming growth factor beta 4 precursor signature

Database

PRINTS; PR01423 TGFBETA

References

MIM; 601877

Literature References 1. KOTHAPALLI, R., BUYUKSAL, I., WU, S.-Q., CHEGINI, N. AND TABIBZADI

Detection of ebaf, a novel human gene of the

transforming growth factor beta superfamily association of gene expre

with endometrial bleeding.

J.CLIN.INVEST. 99 2342-2350 (1997).

2. MENO, C., SAIJOH, Y., FUJII, H., IKEDA, M., YOKOYAMA, T., YOKOYAMA

TOYODA, Y. AND HAMADA, H.

Left-right asymmetric expression of the TGF beta-family member lefty

mouse embryos.

NATURE 381 151-155 (1996).

#### Documentation

Abnormal endometrial bleeding is a common manifestation of gynaecolog diseases. A human gene, termed endometrial bleeding associated factor which is expressed strongly in endometrium, has been shown to be asso with abnormal endometrial bleeding [1]. The predicted protein product ebaf shares similarity with members of TGF-beta superfamily. Thus eba novel member of the TGF-beta superfamily and an endometrial tissue  $f_{\ell}$ whose expression is associated with normal menstrual and abnormal endometrial bleeding [1].

Examples of lateral asymmetry are often found in vertebrates (e.g., t heart being on the left side), but the molecular mechanisms governing establishment of left-right (L-R) handedness are unknown [2]. It is r thought that the gene lefty, a member of the transforming growth fact family, may encode a morphogen for L-R determination [2]. Lefty prote which contains the cysteine-knot motif characteristic of this superfa

is expressed in the left half of gastrulating mouse embryos. This asy expression is transient and occurs just before the first sign of late asymmetry appears [2]. Thus lefty may be involved in establishing L-I asymmetry in the organ systems of mammals [2].

TGFBETA4 is a 14-element fingerprint that provides a signature for the transforming growth factor beta 4 precursor proteins. The fingerprint derived from an initial alignment of 2 sequences: the motifs were dra conserved regions spanning virtually the full alignment length, focus those sections that characterise TGF-beta 4 proteins but distinguish from the rest of the TGF-beta superfamily. Two iterations on SPTR39 1 required to reach convergence, at which point a true set comprising 4 sequences was identified. Several partial matches were also found (Q! Q9W6I7, Q9PW55, Q9W6I6), all antivin or Lefty proteins that match 2-4

# Summary Information

4 codes involving 14 elements 0 codes involving 13 elements 0 codes involving 12 elements O codes involving 11 elements 0 codes involving 10 elements 0 codes involving 9 elements O codes involving 8 elements 0 codes involving 7 elements O codes involving 6 elements 0 codes involving 5 elements 1 codes involving 4 elements 1 codes involving 3 elements 2 codes involving 2 elements

# Composite Feature Index

14	4	4	4	4	4	4	4	4	4	4	4	4	4	4
13	0	0	0	0	0	0	0	0	0	0	0	0	0	0
12	0	0	0	0	0	0	0	0	0	0	0	0	0	0
11	0	0	0	0	0	0	0	0	0	0	0	0	0	0
10	0	0	0	0	0	0	0	0	0	0	0	0	0	0
9	0	0	0	0	0	0	0	0	0	0	0	0	0	0
8	0	0	0	0	0	0	0	0	0	0	0	0	0	0
7	0	0	0	0	0	0	0	0	0	0	0	0	0	0
6	0	0	0	0	0	0	0	0	0	0	0	0	0	0
5	0	0	0	0	0	0	0	0	0	0.	0	0	0	0
4	0	1	0	0	0	0	0	1	0	0	1	1	0	0
3	0	0	0	0	1	0	0	1	0	0	1	0	0	0
2	0	0	2	0	0	0	0	2	0	0	0	0	0	0
	1	2	3	4	5	6	7	8	9	10	11	12	13	14

True Positives	075610	075611	TGF4_HUMAN TG	F4_MOUSE		
True Positive Partials	Q9PVN4	ving 4 elements				
	Q9W6I7	<b>-</b>				
	Codes invol	ving 2 elements Q9W6I6				
Sequence Titles	075610 075611 TGF4 HUMAN TGF4 MOUSE	SIGNALING MOLECUM SIGNALING MOLECUM TRANSFORMING GROW TRANSFORMING GROW	LE LEFTY-A - Homo WTH FACTOR BETA 4	sapiens	s (Huma SOR (TO	an). GF-BETA 4)
	Q9PVN4	LEFTY - Gallus ga	allus (Chicken).			
	Q9W6I7	SIGNALING MOLECU	LE LEFTY2 - Brack	nydanio :	rerio.	(Zebrafish)
	Q9PW55 Q9W6I6	ANTIVIN - Brachyo SIGNALING MOLECU				
Scan History	SPTR39_14f	2 50 NSINGLE		٠		
Initial Motifs		1.1.00				
	Motif 1 wi Element	dth=23	Soon Td	St	Tnt	Pn+
		LAGPGAALTEEQ	Seqn Id TGF4_HUMAN	4	Int 4	Rpt -
		LVSLREALTGEQ	TGF4_MOUSE	4	4	-
•		1.1.00				
	Motif 2 wi Element	dth=22	Soon Td	Q+	Tnt	Dn+
	QILGSLLQQLQ	LDOPPVLDKAD	<b>Seqn Id</b> TGF4 MOUSE	<b>St</b> 26	Int -1	Rpt -
	QLLASLLRQLQ		TGF4_HUMAN	26	-1	-
	Motif 3 wi	dth=17				
	Element		Seqn Id	St	Int	Rpt
	EGMVIPSHVRT	-	TGF4_MOUSE	49	1	-
	EKLVIPAHVRA	QYVVLL	TGF4_HUMAN	49	1	-
	Motif 4 wi	dth=19				
	Element		Seqn Id	St	Int	Rpt
	RSRGKRFSQNL	REVAGRFL	TGF4_MOUSE	72	6	-
	RSRGKRFSQSF	REVAGRFL	TGF4_HUMAN	71	5	-
	Motif 5 wi	dth=16				
	Element		Seqn Id	St	Int	Rpt
	HLLVFGMEQRL		TGF4_MOUSE	97	6	<del>-</del> .
	HLLVFGMEQRL	PPNSE	TGF4_HUMAN	96	6	-
•	Motif 6 wi	dth=21				
	Element		Seqn Id	St	Int	Rpt
	ELVQAVLRLFQ		TGF4_HUMAN	111	-1	-
	ELVQAVLRLFQ	EPVPRTALRR	TGF4_MOUSE	112	-1	-
	Motif 7 wi	dth=19				
	Element		Seqn Id	St	Int	Rpt
	ROKRLSPHSAR		TGF4_MOUSE	132	-1	-
	RHGRLSPAAPK	AKVIVEWL	TGF4_HUMAN	131	-1	-

	Motif 8 width=22				-
	Element	Seqn Id	St	Int	Rpt
	RDDGSNRTALIDSRLVSIHESG	TGF4_MOUSE	153	2	_
	RDDGSNRTSLIDSRLVSVHESG	TGF4_HUMAN	151	1	-
	Motif 9 width=24				
	Element	Segn Id	St	Int	Rpt
	WQQLSRPRQPLLLQVSVQREHLGP	TGF4 MOUSE	187	12	-
	WQQLSRPPEPLLVQVSVQREHLGP	TGF4 HUMAN	185	12	_
		<del></del>			
	Motif 10 width=16				
	Element	Seqn Id	St	Int	Rpt
	HKLVRFAAQGTPDGKG	TGF4_MOUSE	216	5	-
•	HKLVRFASQGAPAGLG	TGF4_HUMAN	214	5	-
	Motif 11 width=26				
	Element	Seqn Id	St	Int	Rpt
	QLELHTLDLKDYGAQGNCDPEAPVTE	TGF4 MOUSE	236	4	-
	QLELHTLDLRDYGAQGDCDPEAPMTE	TGF4 HUMAN	232	2	_
	•		,	_	
	Motif 12 width=22	,			
	Element	Seqn Id	St	Int	Rpt
	RQEMYLDLQGMKWAENWILEPP	TGF4_MOUSE	267	5	-
	RQEMYIDLQGMKWAKNWVLEPP	TGF4_HUMAN	263	5	-
	Motif 13 width=21				
	Element	Seqn Id	St	Int	Rpt
• •	QLPESLTSRWPFLGPRQCVAS	TGF4 MOUSE	301	12	
	QPPEALAFNWPFLGPRQCIAS	TGF4_HUMAN	297	12	_
	Motif 14 width=20				
	Element	Seqn Id	St	Int	Pot
•	PMIVSVKEGGRTRPQVVSLP	TGF4 MOUSE	327	5	Rpt
	PMIVSIKEGGRTRPQVVSLP	TGF4_MOUSE	323	5	_
	THE VOLUME OF THE VALUE OF THE	IGI 4_HOMAN	323	3	_
Final Motifs	Motif 1 width=23				
	Element	Seqn Id	St	Int	Rpt
	LWLCWALWVLPLAGPGAALTEEQ	TGF4 HUMAN .	4	4	-
	LWLCWALWVLPLASPGAALTGEQ	075610	4	4	-
	LWLCWALWALSLVSLREALTGEQ	TGF4 MOUSE	4	4	-
	LWLCWALWVLPLAGPGAALTEEQ	<u>075611</u>	4	4	-
	Motif 2 width=22				
	Element	Seqn Id	St	Int	Rpt
	QILGSLLQQLQLDQPPVLDKAD	TGF4 MOUSE	26	-1	
	QLLASLLRQLQLSEVPVLDRAD	TGF4 HUMAN	26	-1	_
	QLLGSLLRQLQLKEVPTLDRAD	075610	26	-1	_
	QLLGSLLRQLQLSEVPVLDRAD	075611	26	-1	-
	Matif 3 width 17				
	Motif 3 width=17 Element	Com Td	<b>C</b> :	<b>T</b> - •	<b>5</b>
		Seqn Id	St	Int	Rpt
	EGMVIPSHVRTQYVALL EKLVIPAHVRAQYVVLL	TGF4 MOUSE	49	1	-
	EELVIPTHVRAQYVALL	<u>TGF4_HUMAN</u> 075610	49	1	-
	EKLVIPAHVRAQYVVLL		49	1	-
	TYTH A TENTANNÓI A ADD	<u>075611</u>	49	1	-
	Motif 4 width=19				
	Element	Seqn Id	St	Int	Rpt

RSRGKRFSQNLREVAGRFL	TGF4 MOUSE	72	6	_
		71	5	
RSRGKRFSQSFREVAGRFL	TGF4 HUMAN	•		-
RSRGKRFSQSFREVAGRFL	<u>075610</u>	72	6	-
RSRGKRFSQSFREVAGRFL	075611	72	6	-
Motif 5 width=16				
Element	Segn Id	St	Int	Rpt
HLLVFGMEQRLPPNSE	TGF4 MOUSE	97		MPC
			6	-
HLLVFGMEQRLPPNSE	TGF4_HUMAN	96	6	<del>-</del>
HLLVFGMEQRLPPNSE	<u>075611</u>	97	6	-
HLLVFGMEQRLPPNSE	075610	97	6	-
Motif 6 width=21				
Element	Segn Id	St	Int	Rpt
ELVQAVLRLFQEPVPKAALHR	075611	112	-1	
	TGF4 HUMAN		-1	
ELVQAVLRLFQEPVPQGALHR		111		_
ELVQAVLRLFQEPVPKAALHR	075610	112	-1	-
ELVQAVLRLFQEPVPRTALRR	TGF4_MOUSE	112	-1	-
Motif 7 width=19				
Element	Seqn Id	St	Int	Rpt
RQKRLSPHSARARVTIEWL	TGF4 MOUSE	132	-1	-
RHGRLSPAAPKARVTVEWL	TGF4 HUMAN	131	-1	_
				-
RHGRLSPRSAQARVTVEWL	<u>075611</u>	132	-1	-
RHGRLSPRSARARVTVEWL	075610	132	-1	-
Motif 8 width=22				
Element	Segn Id	St	Int	Rpt
RDDGSNRTALIDSRLVSIHESG	TGF4 MOUSE	153	2	
RDDGSNRTSLIDSRLVSVHESG	TGF4 HUMAN	151	1	_
RDDGSNRTSLIDSRLVSVHESG				
	<u>075610</u>	153	2	-
RDDGSNRTSLIDSRLVSVHESG	075611	153	2	-
Motif 9 width=24	•			
Element	Seqn Id	St	Int	Rpt
WQQLSRPRQPLLLQVSVQREHLGP	075610	187	12	
WQQLSRPRQPLLLQVSVQREHLGP	075611	187	12	_
WQQLSRPRQPLLLQVSVQREHLGP	TGF4 MOUSE	187	12	
WQQLSRPPEPLLVQVSVQREHLGP				_
. MÖÖDƏKEREDINÖNƏNÖKEHIĞE	TGF4_HUMAN	185	12	-
Motif 10 width=16				
Element	Seqn Id	St	Int	Rpt
HKLVRFASQGAPAGLG	<u>075610</u>	216	5	-
HKLVRFASQGAPAGLG	075611	216	5	-
HKLVRFAAQGTPDGKG	TGF4 MOUSE	216	5	_
HKLVRFASQGAPAGLG	TGF4 HUMAN	214	5	-
Motif 11 width=26				
Element	Com T-1	<b>^</b> +	<b>T</b> A.	D +
	Seqn Id	St	Int	Rpt
QLELHTLDLRDYGAQGDCDPEAPMTE	075611	234	2	-
QLELHTLDLGDYGAQGDCDPEAPMTE	<u>075610</u>	234	2	-
QLELHTLDLKDYGAQGNCDPEAPVTE	TGF4 MOUSE	236	4	-
QLELHTLDLRDYGAQGDCDPEAPMTE	TGF4 HUMAN	232	2	-
Motif 12 width=22				
Element	Seqn Id	St	Int	Rpt
RQEMYLDLQGMKWAENWILEPP	TGF4 MOUSE	267	5	
RQEMYIDLQGMKWAKNWVLEPP	TGF4 HUMAN		5	_
		263		-
RQEMYIDLQGMKWAENWVLEPP	075610	265	5	-

RQEMYIDLQGMKWAKNWVLEPP	075611	265	5	-
Motif 13 width=21				
Element	Seqn Id	St	Int	Rpt
QLPESLTSRWPFLGPRQCVAS	TGF4 MOUSE	301	12	-
QPPEALAFNWPFLGPRQCIAS	TGF4 HUMAN	297	12	-
QPPEALAFKWPFLGPRQCIAS	<u>0756</u> 10	299	12	-
QPPEALAFNWPFLGPRQCIAS	075611	299	12	-
Motif 14 width=20			•	
Element	Seqn Id	St	Int	Rpt
PMIVSVKEGGRTRPQVVSLP	TGF4 MOUSE	327	5	-
PMIVSIKEGGRTRPQVVSLP	TGF4 HUMAN	323	5	_
PMIVSIKEGGRTRPQVVSLP	075610	325	5	-
PMIVSIKEGGRTRPQVVSLP	075611	325	5	_

# **Hoverprot: TGFB4\_MOUSE**

```
STANDARD;
ID
     TGFB4 MOUSE
                                    PRT:
                                           368 AA.
AC
     Q64280;
DT
     01-NOV-1997 (Rel. 35, Created)
DT
     01-NOV-1997 (Rel. 35, Last sequence update)
     10-MAY-2005 (Rel. 47, Last annotation update)
DT
     Transforming growth factor beta 4 precursor (TGF-beta 4) (Lefty
DΕ
DΕ
     protein) (Lefty-1 protein) (STRA3 protein).
GN
     Name=Ebaf; Synonyms=Lefty, Lefty1, Stra3, Tgfb4;
os
     Mus musculus (Mouse).
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
     Mammalia; Eutheria; Euarchontoglires; Glires; Rodentia; Sciurognathi;
OC
OC
     Muridae; Murinae; Mus.
OX
     NCBI TaxID=10090;
RN
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RP
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RX
     MEDLINE=96202359; PubMed=8610011; DOI=10.1038/381151a0;
RA
     Meno C., Saijoh Y., Fujii H., Ikeda M., Yokoyama T., Yokoyama M.,
RA
     Toyoda Y., Hamada H.;
RT
     "Left-right asymmetric expression of the TGF beta-family member lefty
RT
     in mouse embryos.";
RL
     Nature 381:151-155(1996).
RN
     [2]
     NUCLEOTIDE SEQUENCE.
RP
RA
     Bouillet P.;
     Submitted (JUN-1996) to the EMBL/GenBank/DDBJ databases.
RL
RN
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     NUCLEOTIDE SEQUENCE.
RX
     MEDLINE=98156497; PubMed=9496783;
     Oulad-Abdelghani M., Chazaud C., Bouillet P., Mattei M.-G., Dolle P.,
RΑ
RA
     Chambon P.;
RT
     "Stra3/lefty, a retinoic acid-inducible novel member of the
RT
     transforming growth factor-beta superfamily.";
     Int. J. Dev. Biol. 42:23-32(1998).
RL
RN
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RP
     NUCLEOTIDE SEQUENCE [LARGE SCALE MRNA].
RC
     TISSUE=Embryonic stem cells;
RX
     MEDLINE=22388257; PubMed=12477932; DOI=10.1073/pnas.242603899;
RA
     Strausberg R.L., Feingold E.A., Grouse L.H., Derge J.G.,
     Klausner R.D., Collins F.S., Wagner L., Shenmen C.M., Schuler G.D.,
RA
RA
     Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
     Hopkins R.F., Jordan H., Moore T., Max S.I., Wang J., Hsieh F.,
RA
     Diatchenko L., Marusina K., Farmer A.A., Rubin G.M., Hong L.,
RA
RΑ
     Stapleton M., Soares M.B., Bonaldo M.F., Casavant T.L., Scheetz T.E.,
RA
     Brownstein M.J., Usdin T.B., Toshiyuki S., Carninci P., Prange C.,
RA
     Raha S.S., Loquellano N.A., Peters G.J., Abramson R.D., Mullahy S.J.,
RA
     Bosak S.A., McEwan P.J., McKernan K.J., Malek J.A., Gunaratne P.H.,
RA
     Richards S., Worley K.C., Hale S., Garcia A.M., Gay L.J., Hulyk S.W.,
RA
     Villalon D.K., Muzny D.M., Sodergren E.J., Lu X., Gibbs R.A.,
     Fahey J., Helton E., Ketteman M., Madan A., Rodrigues S., Sanchez A.,
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     Whiting M., Madan A., Young A.C., Shevchenko Y., Bouffard G.G.,
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RA
     Blakesley R.W., Touchman J.W., Green E.D., Dickson M.C.,
RA
     Rodriguez A.C., Grimwood J., Schmutz J., Myers R.M.,
RA
     Butterfield Y.S.N., Krzywinski M.I., Skalska U., Smailus D.E.,
```

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RA
     Schnerch A., Schein J.E., Jones S.J.M., Marra M.A.;
RT
     "Generation and initial analysis of more than 15,000 full-length human
RT
     and mouse cDNA sequences.";
    Proc. Natl. Acad. Sci. U.S.A. 99:16899-16903(2002).
RL
RN
    FUNCTION.
RP
RX
    MEDLINE=98372436; PubMed=9708731; DOI=10.1016/S0092-8674(00)81472-5;
    Meno C., Shimono A., Saijoh Y., Yashiro K., Mochida K., Ohishi S.,
RA
    Noji S., Kondoh H., Hamada H.;
RA
     "Lefty-1 is required for left-right determination as a regulator of
RT
RT
    lefty-2 and nodal.";
RL
    Cell 94:287-297(1998).
CC
     -!- FUNCTION: Required for left-right axis determination as a
CC
        regulator of LEFTY2 and NODAL.
CC
    -!- SUBCELLULAR LOCATION: Secreted.
CC
     -!- DEVELOPMENTAL STAGE: By E8.0, expressed exclusively on the left
CC
        side of developing embryos with expression predominantly in the
CC
        prospective floor plate (PFP). Weak expression in the lateral-
CC
        plate mesoderm (LPM).
CC
    -!- PTM: The processing of the protein may also occur at the second R-
CC
        X-X-R site located at AA 132-135. Processing appears to be
CC
        regulated in a cell-type specific manner.
CC
     -!- SIMILARITY: Belongs to the TGF-beta family.
CC
     ______
    This Swiss-Prot entry is copyright. It is produced through a collaboration
CC
CC
    between the Swiss Institute of Bioinformatics and the EMBL outstation -
CC
    the European Bioinformatics Institute. There are no restrictions on its
CC
    use as long as its content is in no way modified and this statement is not
CC
CC
     ______
    -!- GENE_FAMILY: HBG074429 [ FAMILY / ALN / TREE ]
CC
DR
    EMBL; <u>D83921</u>; BAA12121.1; -; mRNA.
    EMBL; <u>273151</u>; CAA97497.1; -; mRNA.
DR
DR
    EMBL; AJ000082; CAA03909.1; -; mRNA.
DR
    EMBL; AJ000083; CAA03910.1; -; Genomic DNA.
DR
    EMBL; BC050221; AAH50221.1; -; mRNA.
DR
    PIR; S67507; S67507.
DR
    HSSP; P10600; 1TGJ.
DR
    Ensembl; ENSMUSG00000038793; Mus musculus.
DR
    MGI; MGI:107405; Ebaf.
DR
    GO; GO:0005615; C:extracellular space; TAS.
DR
    InterPro; IPR001839; TGFb.
DR
    InterPro; IPR001111; TGFb N.
DR
    Pfam; PF00019; TGF beta; 1.
    Pfam; PF00688; TGFb propeptide; 1.
DR
DR
    PRINTS; PR01427; TGFBETA4.
DR
    ProDom; PD000357; TGFb; 1.
DR
    SMART; SM00204; TGFB; 1.
DR
    PROSITE; PS00250; TGF BETA 1; 1.
    PRODOM; Q64280.
DR
DR
    SWISS-2DPAGE; Q64280.
KW
    Cytokine; Developmental protein; Glycoprotein; Growth factor; Signal.
FT
    DOMAIN
                       74
                                PRODOM: 2002.1: PD037776 11
                 16
FT
    DOMAIN
                 75
                       241
                                PRODOM:2002.1:PD583730
FT
    DOMAIN
                242
                      362
                                PRODOM:2002.1:PD328043 10
FT
    SIGNAL
                1 · 21
                               Potential.
FT
                 22
    PROPEP
                       76
                                Or 135 (Potential).
FT
                77
    CHAIN
                       368
                                Transforming growth factor beta 4.
                158
                       158
FT
    CARBOHYD
                                N-linked (GlcNAc...) (Potential).
FT
    DISULFID
                253
                       266
                                By similarity.
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FT
     DISULFID
                 265
                        318
                                  By similarity.
FT
     DISULFID
                 295
                        353
                                  By similarity.
FT
    DISULFID
                 299
                        355
                                  By similarity.
                368 AA; 41498 MW; 821DAE663C546B5F CRC64;
SQ
     SEQUENCE
     MPFLWLCWAL WALSLVSLRE ALTGEQILGS LLQQLQLDQP PVLDKADVEG MVIPSHVRTQ
     YVALLQHSHA SRSRGKRFSQ NLREVAGRFL VSETSTHLLV FGMEQRLPPN SELVQAVLRL
     FQEPVPRTAL RRQKRLSPHS ARARVTIEWL RFRDDGSNRT ALIDSRLVSI HESGWKAFDV
     TEAVNFWQQL SRPRQPLLLQ VSVQREHLGP GTWSSHKLVR FAAQGTPDGK GQGEPQLELH
     TLDLKDYGAQ GNCDPEAPVT EGTRCCRQEM YLDLQGMKWA ENWILEPPGF LTYECVGSCL
     QLPESLTSRW PFLGPRQCVA SEMTSLPMIV SVKEGGRTRP QVVSLPNMRV QTCSCASDGA
     LIPRRLQP
//
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